

Uranium incorporation in fluorite and exploration of U-Pb dating

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Abstract. The age of ore deposits constitutes a decisive element in understanding their formation. Deciphering their precise chronology may be a challenge in the absence of mineral phases that can be dated by conventional geochronometers. Fluorite is very common either as the major or accessory mineral in a wide variety of ores and may provide information regarding the origin and timing of mineralizing fluid flows. In this contribution, we explore U-Pb dating on fluorite crystals from the world-class carbonate strata-bound fluorite ore of Pierre-Perthuis in Burgundy (Morvan massif, France). Uranium distribution within fluorite is mapped using induced fission-track and synchrotron radiation X-Ray fluorescence nano-imaging, showing that higher uranium content is measured in an overgrowth of fluorite (F_{log}) as a discrete band. Preservation of a micrometer-thick zonation in U, associated with other substituted elements such as Sr, Y, Fe and Zr implies that neither solid-state diffusion nor dissolution-recrystallization did occur. These U-bearing external fluorite overgrowths contain solid inclusions of about 30 μm globular pyrite crystals with a mean $\delta^{34}S$ of $-23.6 \pm 0.4\text{‰}$ V-CDT. We propose that the U incorporation in the fluorite lattice results from the development of a redox front during bacterial sulphate reduction. F_{log} generation sampled and analyzed by LA-ICP-MS on four different crystals provides identical U-Pb ages within the limits of analytical uncertainty. Considered altogether, these four crystals yield an age estimate of 40.0 ± 1.7 Ma, not corrected for matrix-related elemental fractionation. Our results show that fluorite LA-ICP-MS U-Pb geochronology has potential for dating distinct crystal growth stages, although further research should be conducted to evaluate its accuracy.

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1 Introduction

Sedimentary rocks lying unconformably upon a crystalline basement may host large concentrations of authigenic F, Ba, Pb, Zn and U minerals that are of economic interest (Boiron et al., 2002; Gigon et al., 2020; Gigoux et al., 2015; Leach et al., 2005; Sizaret, 2006). These unconformity-related deposits are usually the result of multiple fluid flows inducing mineral dissolution-recrystallization, remobilization and precipitation (Chi et al., 2018; Walter et al., 2018). Knowing the timing of these successive fluid flows is essential for building robust genetic models for mineral deposits. Fluorite, one of the 27

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critical raw materials defined as crucial for the economy of the European Union (European Commission, 2017), is commonly associated in mineral assemblages in veins or strata-bound deposits. Fluorite Sm-Nd geochronology (Galindo et al., 1994; Dill et al., 2011) might help to reconstruct the geological scenario which led to the mineral deposition, together with other direct
35 radiometric dating of authigenic phases, either by bulk dissolution, e.g., Rb-Sr on sphalerite (Nakai et al., 1993), Re-Os on molybdenite (Markey et al., 1998; Stein et al., 2001) and Re-Os on pyrite (Cardon, 2007; Mathur et al., 2000) or by in situ sampling and analysis (e.g., Ar-Ar on adularia (Cathelineau et al., 2012; Mark et al., 2005) Ar-Ar on Mn oxides (Deng and Li, 2017) or U-Pb on uraninite (Alexandre et al., 2009; Martz et al., 2019).

However, it is frequent that fluorite Sm-Nd geochronology produces errorchrons, and additionally, when the age estimation is
40 considered as geologically consistent, results may be unreliable due to poor analytical precision. Moreover, it is sometimes difficult to interpret Sm-Nd ages when they are obtained from the bulk dissolution of a crystal fragment that may be composed of several growth stages. To overcome these issues, a non-radiometric method based on the acquisition of chemical remnant magnetization in fluorite has been developed and employed to date fluorite deposits in areas where the geodynamic evolution and the apparent polar wander path are well documented (Kawasaki and Symons, 2008; Symons, 1994; Symons et al., 2017).
45 Along with other elements, fluorite accepts the incorporation of uranium and lead in its crystal lattice in substitution for Ca^{2+} (typically at low concentrations, from a few ppm to a few %, Piccione et al., 2019). As a result, fission-track (Grønlie et al., 1990) and (U-Th)/He (Evans et al., 2005; Wolff et al., 2015) thermochronology have been tested in fluorite. Wolff et al., (2016) show that He diffusion in fluorite varies depending on elemental substitution in the crystal lattice.

In recent years, in situ U-Pb geochronology has been successfully applied to a growing number of mineral matrices, including
50 calcite and dolomite (Burisch et al., 2017), hematite (Walter et al., 2018), opal (Nuriel et al., 2019), cassiterite (Moscati and Neymark, 2020), Mn-rich chrysocolla and pseudomalachite (Kahou et al., 2020), epidote (Peverelli et al., 2020), nacrite and fluorite (Piccione et al., 2019). Piccione et al., (2019) reported for the first time LA-ICP-MS U-Pb ages acquired on fluorite that are consistent with those of adjacent nacrite crystals, showing that fluorite U-Pb geochronology can constrain the timing of fluid flows, providing favorable U/Pb. However, the incorporation and retention of U and Pb in the crystal structure of
55 CaF_2 are very poorly documented, hence currently limiting the reliability of fluorite U-Pb dating. Moreover, compared to other common lead-bearing minerals, fluorite sampling by laser ablation presents some additional analytical challenges, such as laser-induced damage (De Bonis et al., 2014; Jia et al., 2005; Rafique et al., 2012) and ejection of solid fragments causing variable ablation rates (Gogoll et al., 1996).

The aim of this study is to examine the spatial distribution of uranium and lead and to further explore small-scale LA-ICP-MS
60 U-Pb geochronology in fluorite crystals. The unconformity-related F-Ba ore in Pierre-Perthuis (Morvan massif, France) was chosen as a case study. This ore has been investigated in previous studies (Gigoux et al., 2015, 2016) and a generation of macroscopic geodic fluorite has been dated at 130 ± 15 Ma by Sm-Nd geochronology (Gigoux et al., 2015). As in many other F-Ba deposits, the most commonly used geochronometers are lacking in Pierre-Perthuis, making it an interesting target to gain knowledge on fluorite precipitation ages through fluorite U-Pb dating. Through the use of various petro-geochemical

65 investigations involving multiple analytical methods, including induced fission-track mapping, synchrotron radiation X-Ray
fluorescence nano-imaging, secondary ion mass spectrometry and LA-ICP-MS, this paper will address the following
questions:

- How are U and Pb spatially distributed within micrometer-thick fluorite growth bands?
- How is uranium incorporated in fluorite?
- 70 - Are U and Pb effectively retained in fluorite?
- Can we use fluorite U-Pb geochronology to gain knowledge of the chronology of fluid flows?

2 Geological setting

The studied area is located at the unconformity between the Paris Basin and the French Massif Central (FMC), about 200 km
south east of Paris. The Paris Basin is an intracratonic sedimentary basin that formed above Cadomian and Variscan basement
75 rocks (Guillocheau, 2000). The FMC was structured during the Variscan orogeny (Lardeaux et al., 2014; Vialette, 1973). The
northern edge of the FMC corresponds to the Morvan Massif made up mostly of granitic rocks emplaced during the Early and
Late Carboniferous (Lardeaux et al., 2014; Vialette, 1973). The northern Morvan basement is composed of three granitic units
namely the Avallon, Lormes and Les Settons and a metamorphic unit (Lower Gneiss Unit). The Pierre-Perthuis ore is spatially
related to an unconformity between the Avallon granite basement and the Late Triassic sediments on the northwestern part of
80 the Morvan Massif (Fig. 1a). Two main geological units out crop at Pierre-Perthuis: (1) the Avallon two-mica granite which
shows a weathered zone of about 0.5 to 3 m thickness at its top, and (2) 4 to 8 m of silicified dolostone that corresponds to the
Assise de Chitry Formation (Fm), a Carnian/Norian dolomite formation (235–208 Ma), silicified after the deposition (Fig.
1b). The main fluorite mineralization forms euhedral (cubic habits) fluorite crystals dispersed within the silicified dolomite
rock. A second occurrence forms centimeter cubic fluorite in geodic cavities that crystallized under relatively narrow
85 temperature range of 80-100°C (Gigoux et al., 2016). Both fluorite mineralizations are complemented by two barite stages (an
early small blade-shaped barite stage and a late fan-shaped pink barite stage), three quartz stages (a microcrystalline stage
responsible for the silicification of the Assise de Chitry Fm co-precipitated with chalcopyrite followed by a quartz fringe stage,
and a late granular stage) and at least four sulphide stages.

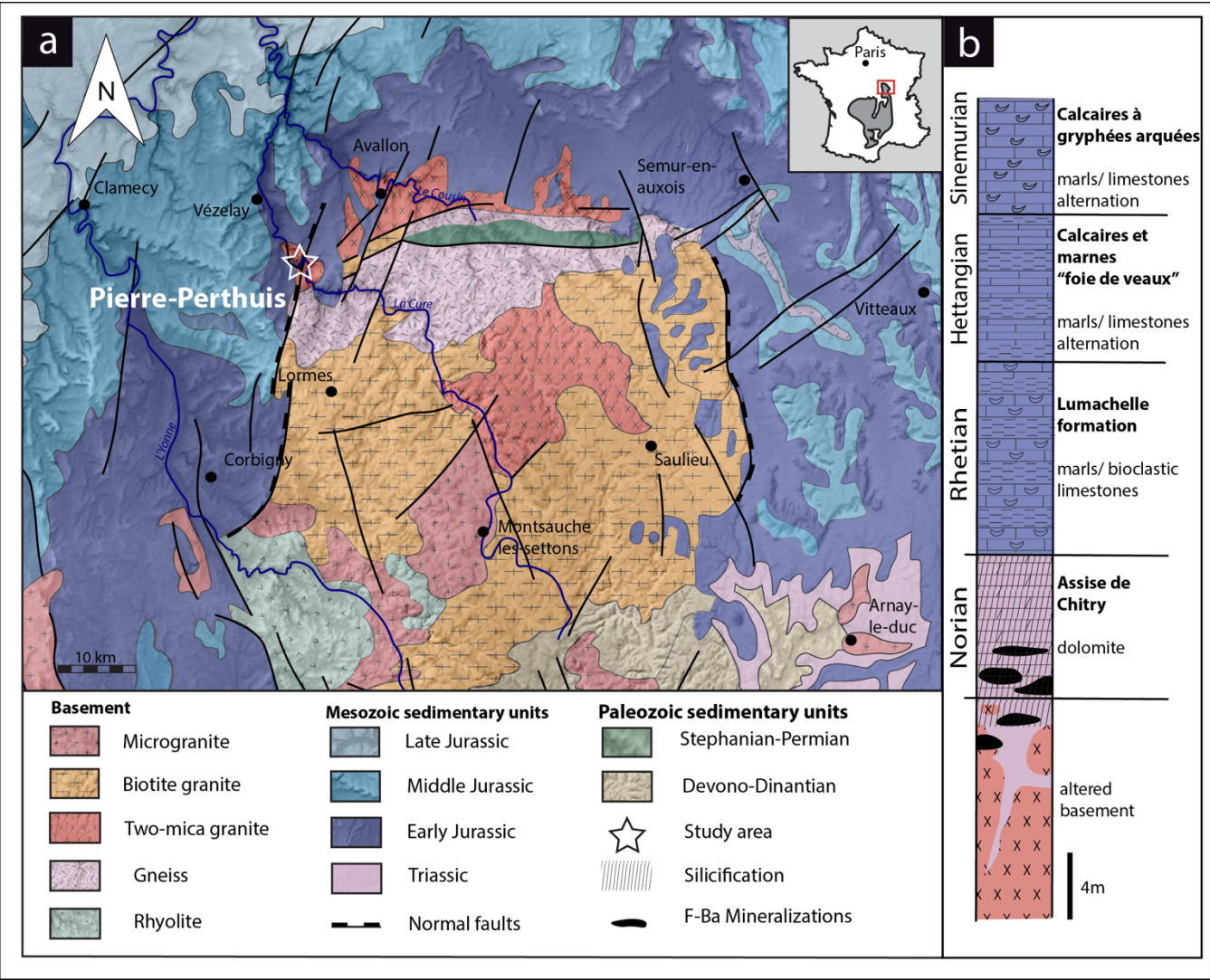
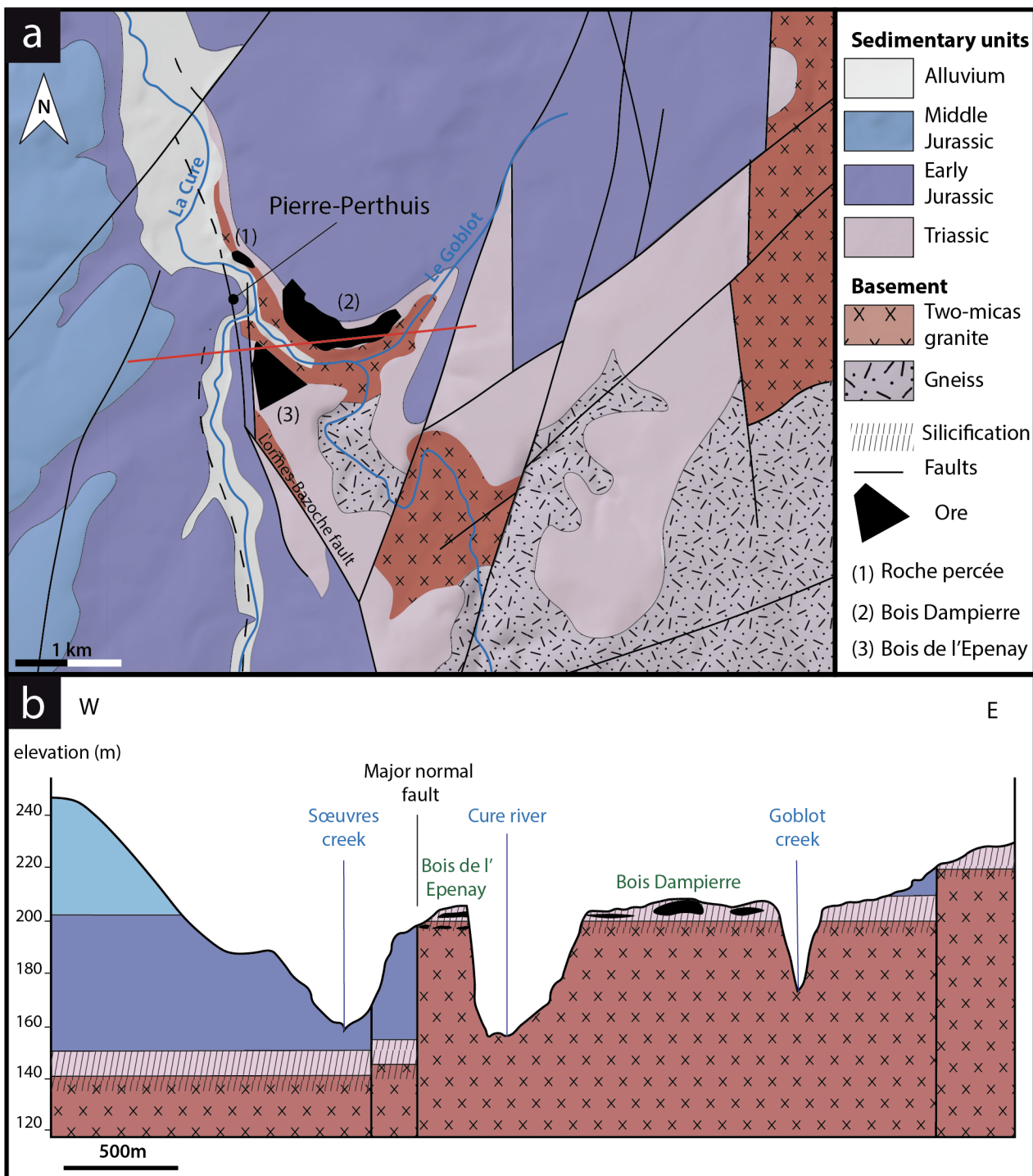


Figure 1: (a) Geological map of the Morvan Massif with the location of the strata-bound fluorite ore of Pierre-Perthuis (modified according to the 1/1.000.000 geological map, BRGM) and its (b) stratigraphic log modified according to Soulé de Lafont and Lhégu (1980).

3 Materials and methods

95 3.1 Sampling strategy

The Pierre-Perthuis district is composed of three areas: (1) Bois Dampierre, (2) Bois de l'Epenay and (3) La roche percée (Fig. 2a). All samples were collected at the Pierre-Perthuis ore from outcropping rocks in the weathered granite (Fig. 2b) close to the sedimentary cover and are summarized in Table 1.



100 Figure 2: a) Location of the mineralized outcrops at Pierre-Perthuis and b) a W-E geological profile localized on the geological map by the red line, modified according to the 1/50.000 geological map of Avallon (Horon et al., 1966 and Gigoux et al., 2016; Soulé de Lafont and Lhégu, 1980).

Sample Name	Location			Applied Method						
	Geographic sector	Lat. X	Long. Y	$\delta^{34}\text{S}$ by SIMS (‰ _{V-CDT})		U-Pb dating by LA-ICP-MS (Ma)		Fission track mapping		Scanning SR-XRF
				$\delta^{34}\text{S}$	2σ	Age	2σ	Tracks nb	Area(μm ²)	
PP18S3	Bois Dampierre	47.43	3.792	−24.2	0.3	N.D.	N.D.	N.D.	N.D.	Zr, Fe, Th, Sr, Pb, Y, U Ca, Fe, Th, Sr, Pb, Y, U
PP18S7	Bois Dampierre	47.43	3.792	−23.6	0.3	N.D.	N.D.	N.D.	N.D.	
PP18S15	Bois Dampierre	47.43	3.792	N.D.	N.D.	N.D.	N.D.	N.D.	N.D.	
PP1802	Bois Dampierre	47.43	3.792	N.D.	N.D.	40.8	3.4	1276	51.5x10 ³	N.D.
PP1801	Roche percée	47.436	3.784	N.D.	N.D.	N.D.	N.D.	196	24x10 ³	N.D.
PPVi2	Roche percée	47.435	3.786	N.D.	N.D.	38.7	1.8	N.D.	N.D.	N.D.
PPVi5	Roche percée	47.435	3.786	N.D.	N.D.	41.1	2.2	N.D.	N.D.	N.D.
PP15	Bois de l'Epenay	47.431	3.786	N.D.	N.D.	41.8	2.8	N.D.	N.D.	N.D.

Table 1: Sample location, sulfur stable isotopic composition of pyrite, fluorite U-Pb age, fission track and Scanning SR-XRF of fluorite. (N.D. = not determined)

3.2 Petrographic observations

Polished thin sections were investigated with optical microscopy in transmitted and reflected light, and under cathodoluminescence (CL) equipment composed of a BX41 microscope coupled to a cold cathode from NewTec operating at 12 kV and 200-300 μA and a Qicam Fast 1394 digital camera. More details about the cathodoluminescence activators and inhibitors in fluorite can be found in Baele et al., (2012) and Kempe et al., (2002).

Uncoated polished thin sections were also observed and analyzed by scanning electron microscopy (SEM, Phenom X Pro) using a charge compensation sample holder in backscattered electron mode (BSE). Images were produced at an acceleration voltage of 15 keV. Semi-quantitative elemental compositions were measured through energy dispersive X-ray spectrometry (EDS) at an acceleration voltage of 15 keV and an accumulation time of 30 s.

3.3 Uranium distribution mapping

To map the uranium distribution in fluorite crystals of the Pierre-Perthuis ore, the fission track cartography based on the induced fission of ²³⁵U was carried out using the external detector method (Gleadow, 1981). Fluorite crystals were mounted in an epoxy resin and polished. Polished sections were covered by muscovite detectors in close contact and irradiated in the Research Neutron Source Heinz Maier-Leibnitz (FRM II) of the Technical University of Munich (Garching, Germany). Thermal neutron fluence was 1 x 10¹⁶ n/cm². Track densities are calibrated using the standard reference glass CN5 with a total U concentration of 12.17 ppm. After irradiation, the external detectors were detached and etched in 48% HF at 20 ± 1°C for 20 minutes. Induced fission tracks revealed on detectors were observed and counted using a LEICA DMLM optical microscope in transmitted light with a x1000 magnification.

According to Enkelmann et al., 2005, the correction factor for uranium concentration between a standard and any unknown geological material depends on the range of latent track lengths and the density of materials. The latent track lengths of fission particles (2R) in fluorite have been simulated using SRIM[®] software (Ziegler et al., 2010). Numerical data employed and results of these simulations are summarized in Table A1. The calculated correction factor (F_{corr}) for uranium concentration in fluorite is 0.866.

3.4 Scanning synchrotron radiation X-Ray fluorescence nano-imaging

Fluorite crystals were mounted in an epoxy resin, polished on both sides to a thickness of a hundred micrometers and cut into 5mm edge squares. These mounts were fixed on Teflon rings and positioned perpendicularly to the incoming X-ray beam.

Seven fluorite samples were scanned by synchrotron radiation X-ray fluorescence (SR-XRF) using the Nanoscopium hard X-ray nanoprobe beamline (Bergamaschi et al., 2017; Somogyi et al., 2015) of synchrotron SOLEIL (Université Paris-Saclay, Saint Aubin, France). Elemental distribution mapping was acquired from these polished sections in different areas at a spatial resolution varying from 50 nm to 1 μm and with an integration time per pixel varying from 150 ms to 20 ms. The 18 keV monochromatic X-ray beam was focused on the crystal samples by a Kirkpatrick-Baez nano-focusing mirror. This excitation energy of 18 keV was chosen farther away from the U-L₃ absorption edge (17.168 keV) in order to avoid the overlapping of the U-L₃ X-ray lines and the Compton scattering peak of the incident X-ray beam from the sample. This allows the optimization of the analytical sensitivity of U due to the reduced spectral background below the U-L₃ lines. To obtain nanometer-scale resolution elemental maps on mm² sized samples, a fast-continuous scanning technique (FLYSCAN) has been employed (Lanzirotti et al., 2010; Medjoubi et al., 2013). Full XRF spectra were collected at each pixel of the scans by two silicon drift detectors of 50 mm² useful area (KETEK H50, KETEK GmbH) used with XMAP (XIA LLC) fast digital multichannel analyzer cards. The XRF detectors were placed at 20° relative to the sample surface (Somogyi et al., 2015).

The high-resolution scans consisted of 100 000 - 300 000 individual XRF spectra corresponding to an XRF spectrum at each image pixel. In order to treat this data-set, as a first step we used the XRF sum-spectrum of the entire scanned area to identify the main elements present. The distribution maps of these elements were reconstructed by an in-house Matlab[®] code. Because the scanned areas sometimes included different mineral matrices (fluorite and pyrite), or successive growth bands of distinct chemical composition in the same mineral, sub-zones were defined by the Matlab[®] code from the elemental distribution maps in order to extract the average XRF spectrum corresponding exclusively to the targeted mineral matrix or growth band. These sub-zones, or regions of interest (ROIs), were selected by using ImageJ[®] software (Schneider et al., 2012). The mean XRF spectra of the ROIs was fitted using PyMCA[®] software (Solé et al., 2007). This allowed us to identify additional elements, present only in tiny areas or in small quantities (e.g., Th), and as such not detectable in the sum-spectrum of the whole sample area. As a second step, the elemental distribution maps of all the identified elements were reconstructed by correcting for spectral overlaps depending on the intensity ratios of the X-ray lines. Because sample thickness varies between 100 to 300

micrometers, the low energy X-ray spectral line intensities will not be revealed from the whole sample thickness by SR-XRF.
155 The depth of information for all elements included in the present paper is documented in Table A3.

3.5 Sulfur stable isotope composition of pyrite

Sulfur isotopes were measured by Secondary Ion Mass Spectrometry (SIMS) using a multi-collectors CAMECA IMS1270 E7 microprobe at the *Centre de Recherches Pétrographiques et Géochimiques* (CRPG, Université de Lorraine and CNRS) in Nancy (France). Samples that were already prepared for synchrotron XRF-spectromicroscopy were re-used. They were glued
160 on a glass slide, then coated with a layer of 30 nm of Au. To accurately locate the SIMS analysis, petrographic observations using reflected light microscopy were carried out before and after gold plating. Pyrite inclusions in fluorite were sputtered with a 10 kV Cs⁺ primary ion beam with a current of 2.7 nA. A normal-incidence electron gun was used for charge compensation. ³²S⁻ and ³⁴S⁻ secondary ions were detected in multi-collection mode using two Faraday cups. Samples were pre-sputtered for 120 seconds with a beam raster of 20 x 20 μm prior to signal acquisition to minimize surface contamination.
165 Faraday cup backgrounds were measured during the pre-sputtering before each analysis and then used for correcting the data. Typical count rates of ³²S⁻ and ³⁴S⁻ were of 4.8×10⁸ counts per second (cps) and 2×10⁷ cps respectively, during 30 cycles of 4.96 seconds acquisition time. The spot size was about 15 μm in diameter. A nominal mass resolution (M/ΔM) of 4998 was used to resolve interference with hydrides.

A Sierra Pyrite sample was used as an internal matrix-match standard to correct the sulfur isotopic ratio. Data was integrated
170 as a ³²S/³⁴S ratio (δ³⁴S in ‰) and referenced to the Vienna-Canyon Diablo Troilite (V-CDT) scale (Ding et al., 2001). Homogeneity of the Sierra Pyrite has been documented by LaFlamme et al., (2016) by three analytical techniques (EPMA: Electron Probe Micro Analyser, Wavelength-Dispersive X-Ray spectroscopy-maps and SEM-BSE) and its δ³⁴S measured by fluorination gas-source mass spectrometry at 2.17 ± 0.08 ‰_{V-CDT}. Measurements were carried out through daily sequences consisting of 5 analyses of matrix-match reference material followed by spots on unknown samples. For the first
175 session, Sierra pyrite was measured at 0.94 ± 0.02 ‰_{V-CDT}, and at 0.92 ± 0.02 ‰_{V-CDT} for the second analytical session. Instrumental mass fractionation (IMF) for ³⁴S/³²S was quantified during each analytical session using all the replicated analyses of the matrix-match standard. During the first session the IMF was -1.23 ± 0.34 ‰_{V-CDT}, and -1.25 ± 0.28 ‰_{V-CDT} for the second. Final uncertainties are reported at a 95% confidence level (2σ), propagating the systematic uncertainty of primary reference material Sierra pyrite (3.7 ‰, LaFlamme et al., 2016), and the propagated correction for instrument mass
180 fractionation through the analytical session by quadratic addition.

3.6 In situ U-Pb geochronology

Uranium and lead isotopic compositions were measured for 4 fluorite samples showing fluorite overgrowth (F_{log}): (1) PPVi2 (n = 16), (2) PPVi5 (n = 20), (3) PP1802 (n = 30) and (4) PP15 (n=26), using a 193 nm excimer laser (Teledyne, Photon Machines) coupled to a Thermo ScientificTM Sector Field Inductively Coupled Plasma Mass Spectrometer

185 (SF-ICP-MS) ELEMENT XR (ThermoFisher Scientific, Waltham, USA) at the Geosciences laboratory (GEOPS) of the University Paris-Saclay.

The overall analytical procedure is derived from LA-ICP-MS U-Pb geochronology of carbonates (Roberts et al., 2020), developed at the University Paris-Saclay for calcite (Brigaud et al., 2020) - see Table A4 for analytical details. The main difference concerns the laser energy needed to ablate fluorite. Indeed, a fluence of 6 J.cm^{-2} was required to ablate our
190 natural fluorite crystals. To properly correct for downhole fractionation (DF) and other sources of laser-induced elemental fractionation (LIEF) on the measured $^{206}\text{Pb}/^{238}\text{U}$ ratios, a fluorite primary reference material (RM) should be used. However, such a fluorite RM does not exist yet. Thus, similarly to Piccione et al., (2019), we used a calcite RM, the Permian Reef Complex Walnut Canyon WC-1 (Roberts et al., 2017) as the primary reference material. We highlight that our fluorite U-Pb ages are thus not corrected for matrix-related elemental fractionation. To evaluate the accuracy of our U-Pb ages, we included
195 the analysis of a fluorite sample independently dated by (U–Th–Sm)/He thermochronology (Wolff et al., 2016). It corresponds to a sodium-and Rare Earth Element (REE)-rich green fluorite from the Horni Krupka deposit in Czech Republic (named “HK13”) that has been dated by Wolff et al., (2016) at $290 \pm 10 \text{ Ma}$.

The laser beam diameter for fluorite unknowns was $85 \text{ }\mu\text{m}$. Fluorite crystals were ablated at a repetition rate of 10 Hz and a fluence of 6.25 J.cm^{-2} . Calcite reference materials, including WC-1 primary standard and two secondary standards that are
200 detailed below, were ablated at a repetition rate of 8 Hz and a fluence of 1 J.cm^{-2} , and a beam diameter of $150 \text{ }\mu\text{m}$. Glass reference materials NIST612 and NIST614 (Jochum et al., 2011) were ablated at a repetition rate of 10 Hz, a fluence of 6.25 J.cm^{-2} and a beam size of $40 \text{ }\mu\text{m}$ for NIST612 and $110 \text{ }\mu\text{m}$ for NIST614.

Each analysis consists of 30 s background acquisition followed by 30 s of sample ablation and 30 s washout. Prior to analysis, each spot was pre-ablated for 7 s at a frequency of 10 Hz and with a fluence of 6.25 J.cm^{-2} for reference material (NIST) and
205 fluorite samples, and at a frequency of 8 Hz and with a fluence of 1 J.cm^{-2} for calcite reference materials. These pre-ablations are over an area larger than the beam diameter to clean the surface ($155 \text{ }\mu\text{m}$ for calcite, $50 \text{ }\mu\text{m}$ for NIST612, $110 \text{ }\mu\text{m}$ for fluorite and $135 \text{ }\mu\text{m}$ for NIST614) and remove potential surficial Pb contamination. The laser-induced aerosol was carried by helium (He: large volume at 0.5 l.min^{-1} and inner cup at 0.375 l.min^{-1}) from the sample cell to a mixing funnel in which the sample and He are mixed with 0.950 to 1 l.min^{-1} argon to stabilize the aerosol input to the plasma. Signal strength of the ICP-MS was
210 tuned for maximum sensitivity while keeping Th/U at 1.02 and ThO/Th below 0.3% on NIST612. Isotopes ^{206}Pb , ^{207}Pb , ^{208}Pb , ^{232}Th and ^{238}U were acquired with integration times per peak of 10 ms for ^{208}Pb , ^{232}Th , ^{238}U , of 30 ms for ^{206}Pb and of 40 ms for ^{207}Pb by 70 runs.

We used NIST614 to correct for $^{207}\text{Pb}/^{206}\text{Pb}$ fractionation (Jochum et al., 2011). For mass-bias correction of the measured $^{238}\text{U}/^{206}\text{Pb}$ ratios, we used the WC-1 calcite reference material (Roberts et al., 2017). Two secondary calcite reference
215 materials were analyzed during the analytical sessions: Duff Brown Tank (DBT), dated at $64.0 \pm 0.7 \text{ Ma}$ by U-Pb isotope dilution – ICP-MS (Hill et al., 2016), and AUG-B6, a calcite breccia dated by LA-ICP-MS U-Pb at $42.99 \pm 1 \text{ Ma}$ (Pagel et al.,

2018). Internal secondary reference calcite AUG-B6 comes from the Gondrecourt graben (Eastern Paris Basin), part of the European Cenozoic Rift System and has been routinely analyzed for two years at GEOPS (University Paris-Saclay). Measurements have been made by sequences starting with 6 reference material analyses (2 NIST612, 2 NIST614 and 2 WC-1), then a repetition of 10 spots on unknown fluorite followed by 8 reference material analyses (2 NIST614, 2 WC-1, 2 DBT and 2 AUG-B6), 10 spots on unknown fluorite, etc., and ending with 6 reference material analyses (2 NIST612, 2 NIST614 and 2 WC-1). Data was acquired in fully automated mode overnight in two sequences of 336 analyses during about 12 hours of analysis the 19 and 20 December 2019.

Data was reduced in Iolite© using the NIST614 glass as the primary reference material to correct for baseline, for Pb isotope mass bias and for $^{206}\text{Pb}/^{238}\text{U}$ instrumental drift over the sequence time (Lawson et al., 2018; Paton et al., 2011). No down-hole fractionation correction is applied in Iolite© (Nuriel et al., 2017). The two-sigma errors in $^{207}\text{Pb}/^{206}\text{Pb}$ and $^{206}\text{Pb}/^{238}\text{U}$ ratios measured on NIST614 during the analytical session were propagated to the final age uncertainty of fluorite samples by quadratic addition. During the first session, NIST614 two sigma error on $^{207}\text{Pb}/^{206}\text{Pb}$ was 0.31 % and 1.10 % for $^{206}\text{Pb}/^{238}\text{U}$ and for the second session the two-sigma errors on $^{207}\text{Pb}/^{206}\text{Pb}$ and $^{238}\text{U}/^{206}\text{Pb}$ were 2.20% and 0.69% respectively. A signal peak of ^{206}Pb , ^{207}Pb or ^{238}U can occur due to unstable ablation conditions, causing an unusually high $^{206}\text{Pb}/^{238}\text{U}$ or $^{207}\text{Pb}/^{206}\text{Pb}$ standard error (2σ), typically greater than 20 %. In Figure A1, we document four examples of LA-ICP-MS intensity signals during fluorite ablation together with the corresponding 3D surface images of laser craters acquired by optical profilometry. Thirty-five spots, having 2σ errors greater than 20 %, were excluded (5 on PPVi5, 7 on PPVi2, 15 on PP1802 and 8 on PP15). Each reduced piece of data is plotted in a $^{238}\text{U}/^{206}\text{Pb}$ versus $^{207}\text{Pb}/^{206}\text{Pb}$ Tera-Wasserburg graph using IsoplotR (Vermeesch, 2018). An isochron is drawn and the isochron age is deduced by the intersection on the concordia. For each sequence, the age and uncertainty of WC-1 reference calcite, following normalization using NIST614 glass, was calculated without further normalization using a Tera-Wasserburg intercept age. Following the analytic run, we applied a linear correction factor to correct the $^{206}\text{Pb}/^{238}\text{U}$ so that the primary WC-1 yields the correct intercept age (254.4 ± 6.4 Ma, Roberts et al., 2017). For the first session, we obtained 167.0 ± 3.7 Ma for WC-1 with 23 analyses, hence we applied a linear correction factor of 0.656 to correct all $^{206}\text{Pb}/^{238}\text{U}$ ratios of secondary calcite reference materials and fluorite unknowns. For the second analytical session, WC-1 was dated at 148.0 ± 2.3 Ma with 39 analyses. Then, the linear correction factor on the $^{206}\text{Pb}/^{238}\text{U}$ ratio was 0.582. We fixed the $^{207}\text{Pb}/^{206}\text{Pb}$ ratio at 0.85 for the common Pb based on Stacey and Kramers (1975), when we calculated the Tera-Wasserburg intercept age on WC-1 (Roberts et al., 2017). Regarding unknown fluorite samples, Tera-Wasserburg intercept ages are calculated by plotting each spot from a single sample and by applying the linear correction factor found on WC-1 for the corresponding session to correct the $^{206}\text{Pb}/^{238}\text{U}$ ratios. An age is then calculated without fixing the initial $^{207}\text{Pb}/^{206}\text{Pb}$ value. Error ellipses of each spots and the error on the Tera-Wasserburg intercept age are 2σ . In each Tera-Wasserburg graph, a first age uncertainty is given that does not include uncertainty propagations (except uncertainties related to the decay constants of ^{235}U and ^{238}U). A second age uncertainty in square brackets is given, by propagating the systematic uncertainty of primary reference material WC-1 age (2.6 %, Roberts et al., 2017) and the two-sigma error of the $^{207}\text{Pb}/^{206}\text{Pb}$

250 and $^{206}\text{Pb}/^{238}\text{U}$ of the analytical session by quadratic addition. Calculated ages for calcite secondary reference materials analyzed during the two sequences were 40.7 ± 1.6 Ma and 44.4 ± 2.0 Ma for AUG-B6 and 63.7 ± 2.2 Ma and 64.6 ± 2.9 Ma for DBT, without fixing the initial $^{207}\text{Pb}/^{206}\text{Pb}$ ratio. These ages are identical within analytical uncertainty to the ages published for these two calcites (Pagel et al., 2018; Hill et al., 2016, respectively). “HK13” fluorite was dated at 285.9 ± 29.3 [30.9] Ma. Even though the isochron is less well defined, this U-Pb age matches the (U-Th-Sm)/He age obtained by Wolff et al.,
255 (2016) at 290 ± 10 Ma.

Tera-Wasserburg diagrams of primary and secondary calcite reference materials and metadata for LA-ICP-MS U-Pb ages of fluorite can be found in Fig. A2 and A3 and Table A4 of Appendix A.

4 Results

4.1 Petrographic observations

260 Three fluorite generations are described below and illustrated in Fig. 3a.

The first fluorite stage is composed of $50\text{ }\mu\text{m}$ up to a millimeter-large white cubes disseminated in the altered granite and mainly in the Assise de Chitry Fm, named euhedral fluorite Fl_{euh} (Gigoux et al., 2016). These crystals may sometimes form centimeter-sized aggregates (Fig. 3a).

The second stage fills pluri-centimetric to pluri-decimetric-large geodic cavities or centimetric veinlets. This geodic fluorite
265 (Fl_{geo}) forms millimetric to centimetric well-developed white and yellow translucent cubes which are more or less interlinked (Fig. 3b). These crystals are fractured and contain many fluid inclusions (Fig. 3c).

Geodic fluorites (Fl_{geo}) sometimes display one or several overgrowths (Fl_{og}). These overgrowths form the third and last stage of fluorite. They appear translucent in transmitted plane polarized light and do not contain any fluid inclusions (see Fig. 3c and Fig. A4). From the center to the border, two growth phases have been noted (1) a translucent fluorite initiated by one or
270 several purple rims (Fl_{og1}) (2) a second growth stage of fluorite with a laminar texture (Fl_{og2}), delineated by a fringe of well-aligned small globular pyrite crystals of about $30\text{ }\mu\text{m}$ (Fig. 3e and 3f). These pyrite crystals are rarely oxidized along fractures (Fig. 3d).

Fission track mapping carried out on Fl_{geo} and $\text{Fl}_{\text{og1\&2}}$ shows that Fl_{geo} and Fl_{og1} are almost devoid of tracks, while a high track density is observed in Fl_{og2} (Fig. 3g and 3h). Uranium is only detected and measured in Fl_{og2} , with U concentration
275 varying from 4 to 50 ppm according to the track counting zones with an average from 7 to 26 ppm per sample (see Table A2 for details).

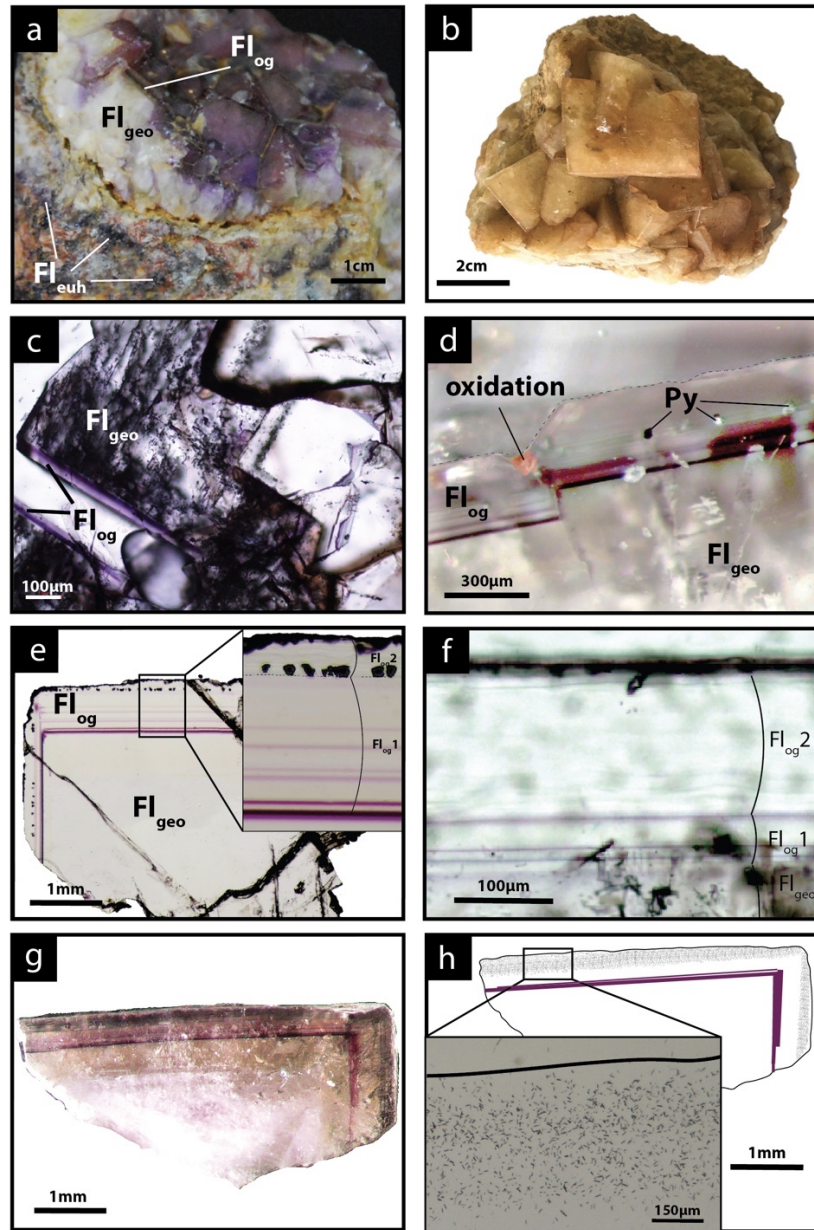


Figure 3: Photographs of the fluorite stages from the Pierre-Perthuis ore. (a) Geode in the altered granite with milky white cubic crystals (Fl_{geo}) and the translucent fluorite overgrowth (Fl_{og}) delimited by a thin purple band. The euhedral fluorite (Fl_{euh}) forms patches in the Avallon granite, from Gigoux et al., 2015, (b) Centimetric cubes of Fl_{geo} , (c) Transmitted light microscopy photograph of Fl_{geo} and the overgrowth corresponding to Fl_{og} , (d) Photograph of the limit between Fl_{geo} and Fl_{og} with a pyrite crystal affected by a fracture and oxidized, (e) Transmitted light microphotograph of a geodic fluorite crystal with the two stages constituting Fl_{og} (1) a translucent fluorite initiated by one or several purple rims: Fl_{og1} ; (2) a final growth stage of laminar fibrous fluorite with pyrite inclusions: Fl_{og2} , (f) Microphotograph in transmitted light focused on Fl_{og2} showing a laminar texture (g) Photograph in transmitted light of an irradiated crystal of $Fl_{geo} + Fl_{og}$, (h) Corresponding interpreted map of the induced fission tracks distribution in a crystal illustrated in (g) with a zooming on the muscovite detector.

4.2 Sulfur stable isotope composition

SIMS analyses were carried out for two fluorite crystals containing pyrite inclusions in their rims (Fig. 4). Five pyrite crystals in PP18S3 sample have $\delta^{34}\text{S}$ values varying from -23.1 to -24.9 ‰_{V-CDT}. The sulfur isotopic ratio of pyrite from PP18S7 sample varies from -19.9 to -24.7 ‰_{V-CDT}. The mean of $\delta^{34}\text{S}$ values for PP18S3 and PP18S7 are -24.2 and -23.6 ‰_{V-CDT}, respectively. Two-sigma uncertainty is below 0.4 ‰_{V-CDT}.

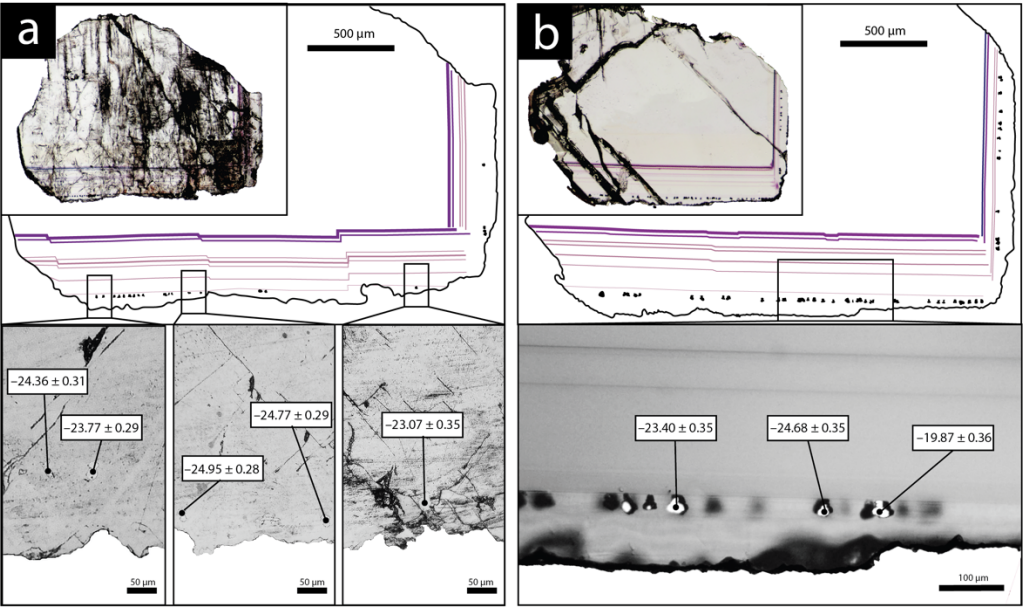


Figure 4: Location of SIMS analyses and the $\delta^{34}\text{S}$ measurements presented in ‰_{V-CDT} for (a) sample PP18S3 and (b) sample PP18S7. Micro-photographs in upper half are in transmitted plane polarized light and microphotographs in the bottom are gold metallized zoomed areas in reflected plane polarized light.

4.3 SR-XRF mapping

We studied in detail two samples that contain the last two generations of fluorite (Fl_{geo} and Fl_{log} described above) to investigate geochemical heterogeneities. PP18S7 shows several pyrite crystals at various depths while PP18S15 shows a single pyrite at the surface of the sample (Fig. 5a and 5b respectively). Synchrotron radiation XRF mapping of a transect from Fl_{geo} to Fl_{log2} shows areas of lower concentration of Ca within the Fl_{log2} exclusively (Fig. 5b), and higher concentrations of substituted elements such as Zr, Y, Sr, U, Th and Pb (Fig. 5a). Lead is present both into pyrite crystals (Fig. 5a) and in Fl_{log2} (Fig. 6).

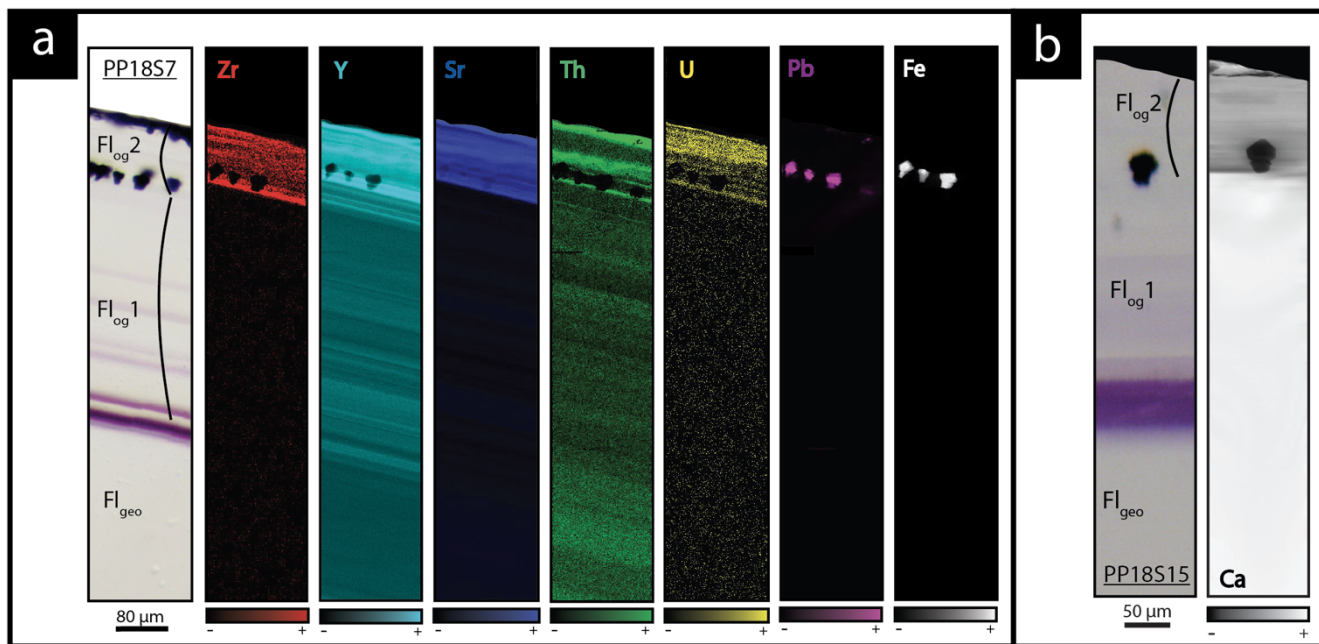
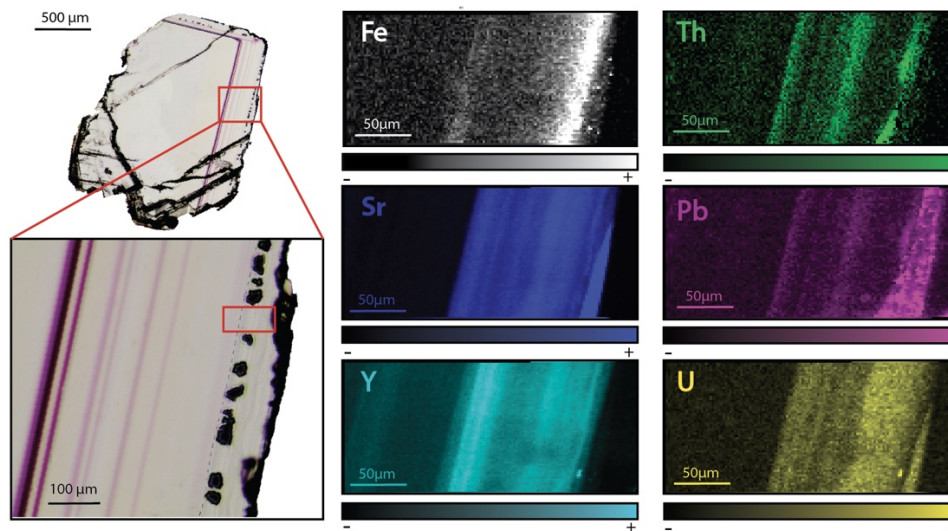


Figure 5: Transmitted plane polarized light microphotographs and corresponding synchrotron radiation X-ray fluorescence elemental maps through a transect from the core crystal of Fl_{geo} to the Fl_{og} rim from, a) sample PP18S7 and b) PP18S15. Black areas indicate concentration below the limit of detection of the element. Globular pyrite crystals are included in Fl_{og2} .

Other maps have been generated with a focus on the transition between Fl_{og1} and Fl_{og2} (Fig. 6), highlighted as distinct by petrographic observations and fission track mapping (Fig. 3f and 3h). This map reveals elemental substitution and uranium incorporation. At this scale, SR-XRF mapping shows enrichment in Fe, Th, Sr, Pb, Y and U in Fl_{og2} . Lead was detected by SR-XRF mapping, and its spatial distribution in Fl_{og2} is correlated with thorium (Fig. 6) but not with uranium (Fig. A5).



315 **Figure 6:** SR-XRF imaging of the transition between Fl_{og1} and Fl_{og2} in PP18S15 showing the distribution of Fe, Th, Sr, Pb, Y and U contents. Map dimensions are 261μm x 122μm.

A focus on a pyrite crystal in Fl_{og2} reveals concentric zonation in Cu, As and Ni concentrations (Fig. 7).

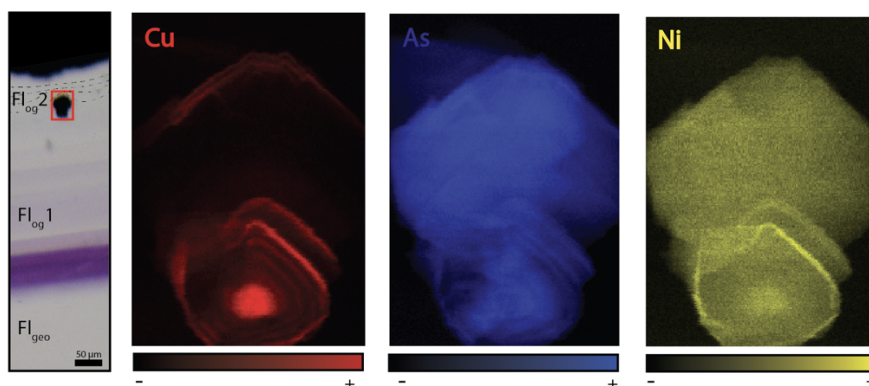


Figure 7: Copper, arsenic and nickel distribution maps of a pyrite included in Fl_{og2}. Maps dimensions are 52 μm x 76 μm.

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4.4 Laser ablation U-Pb geochronology

Four isochrons were calculated, with a common initial $^{207}\text{Pb}/^{206}\text{Pb}$ composition ranging from 0.815 to 0.840 (Tera-Wasserburg graphs were generated without anchoring the upper intercept of the discordia line to a given $^{207}\text{Pb}/^{206}\text{Pb}$ composition). Outer rims of geodic fluorite crystals from Pierre-Perthuis yield an age of 38.7 ± 1.5 [1.8] Ma with 9 analyses and a MSWD of 0.68 (Fig. 8a) for PPVi2, 41.1 ± 1.8 [2.2] Ma (MSWD = 1 and n=15) for PPVi5 (Fig. 8b), 40.8 ± 3.0 [3.4] Ma for PP1802 with 11 analyses and a MSWD = 0.6 (Fig. 8c) and PP15 of 41.8 ± 2.4 [2.8] Ma (MSWD=1.5 and n=22, Figure 8d). As these ages are identical within analytical uncertainties, all data from the different crystals but corresponding to the same generation according

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to petrography, were then plotted in a single graph (Fig. 8e), giving an age of 40.0 ± 0.9 [1.7] Ma (MSWD=1.2). Because U is variable in the samples, data is well distributed in the Tera-Wasserburg space, and the isochron is rather well constrained.

330 To evaluate the accuracy of fluorite LA-ICP-MS U-Pb dating in the absence of calibration against a fluorite primary RM, we tested the HK13 fluorite, a sample independently dated at 290 ± 10 Ma by (U–Th–Sm)/He thermochronology (Wolff et al., 2016). Although we found an age of 285.9 ± 30.9 Ma (Fig. A3), interpretation of results is somehow limited by a poor data spread in a Tera-Wasserburg diagram and by rather large uncertainties regarding single analytical spots related to unstable ablation rates.

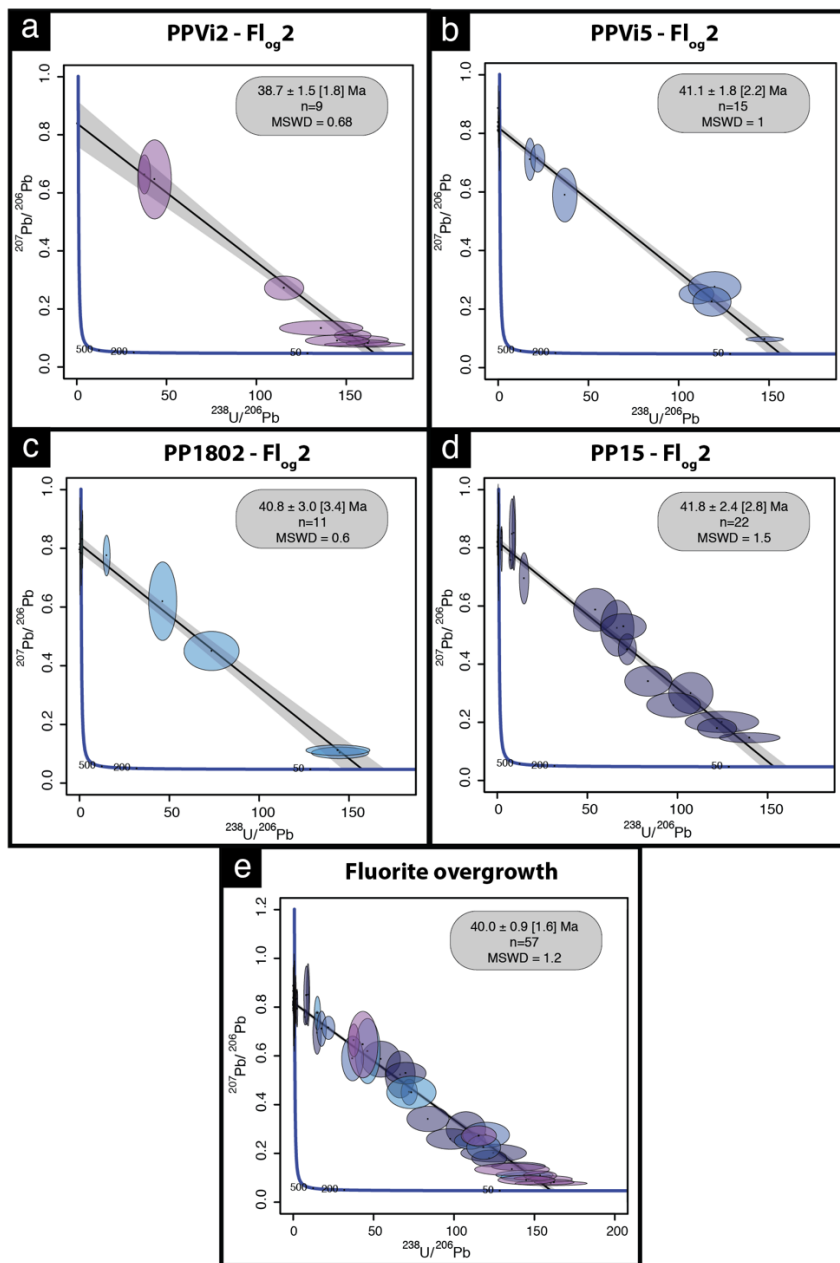


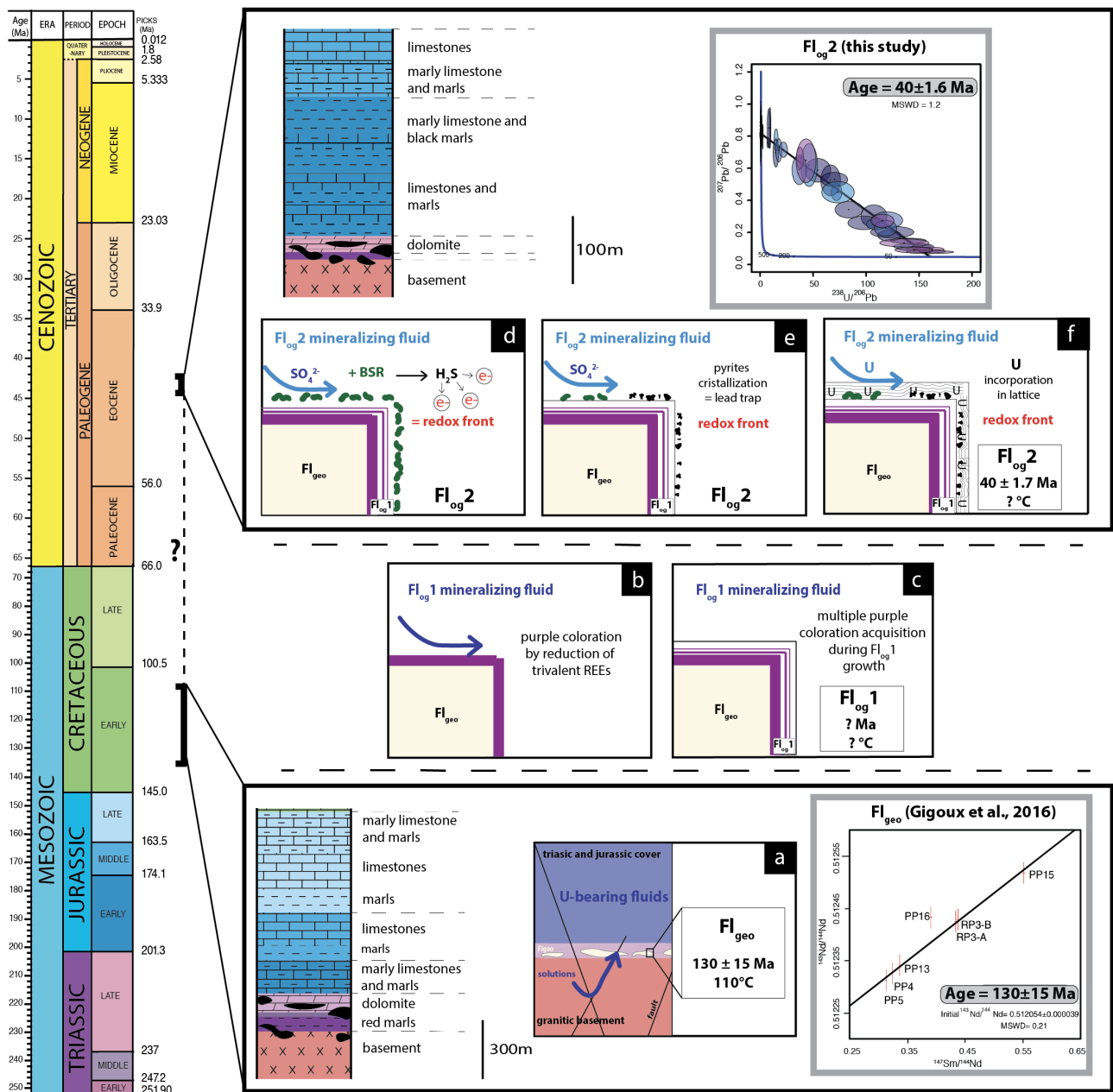
Figure 8: Tera-Wasserburg for fluorite. (a,b,c and d) U-Pb ages of the four fluorite overgrowth (FI_{og2}) samples. (e) Isochron obtained from the compilation of all data illustrated in a,b,c and d. “n” is the number of analytical spots for each sample.

5 Discussion

5.1 Uranium and lead incorporation in fluorite

340 The fluorite overgrowths are composed of two distinct layers: (1) Flog1 that is devoid of U and (2) Flog2, in which U was effectively incorporated within the crystal and correlated with precipitation of pyrite. Flog2 appears colorless with an internal fibrous texture. We did not observe any pleochroic halos like Dill and Weber (2010) have documented in fluorite around actinide-rich inclusions. Fission track mapping instead shows a homogeneous incorporation of U (Fig. 3g and 3h). On a smaller scale, SR-XRF reveals micrometer-scale fluctuations in uranium concentrations along the growth directions, correlated with
 345 the incorporation of other substituted elements such as Sr and Zr (Fig. 5). The incorporation of Zr in the fluorite lattice, together with U, may reflect a contribution from the leaching of volcanic glasses. Indeed, correlations between uranium and zirconium concentrations have been documented in various U deposits spatially associated to volcanic rocks or volcanic fragments reworked in sandstones and conglomerates (Forbes et al., 1984; Cinelu and Cuney, 2006; Pons, 2015). Late Variscan ignimbrites are cropping out ca. 35 km south of Pierre-Perthuis (Carpena et al., 1984).

350 The transition between Flog1 and Flog2 may be explained by the development of a redox front, causing uranium incorporation in the crystal lattice. The presence of numerous pyrite crystals in Flog2, similar to those reported by Dill and Weber (2010), characterizes a reducing environment that has probably initiated the precipitation of U. As an analogy, it can be mentioned that pyrite is common in roll-front uranium deposits (Bonnetti et al., 2015; Cai et al., 2007; Campbell et al., 2012; Ingham et al., 2014). Globular pyrites, similar to those observed in Flog2, are usually interpreted as biogenic in origin (Belyi et al., 1972; Blakeman et al., 2002; Chen et al., 2019; Bonnetti et al., 2020; Cai et al., 2007; Rackley, 1972). This hypothesis is
 355 supported by the sulfur stable isotope composition of the pyrites included in Flog2 (mean $\delta^{34}\text{S} = -23.6 \text{ ‰}_{\text{V-CDT}}$). The homogeneous $\delta^{34}\text{S}$ of these pyrites indicates a large amount of sulfate available for reduction by bacteria in an open system (Hough et al., 2019; Magnall et al., 2016). Concentric incorporation of Ni, As, and Cu (Fig. 7) were reported both in biogenic (Baumgartner et al., 2020) and abiogenic pyrite (Wu et al., 2019). Barbarand et al., (2013) show that the basement of the
 360 southeastern Paris Basin was subjected to temperatures less than 60 °C during the Eocene period. This temperature estimate is compatible with bacterial activity. Hence, we propose that uranium incorporation in Flog2 is related to bacterial sulphate reduction (BSR, Fig. 9). The solubility of uranium was enhanced by F-bearing solutions (Xing et al., 2019) that leached the granitic basement through the fracture network (Fig. 9a). When reaching the unconformity between the basement and the sedimentary cover, a local redox front generated by BSR causes U to precipitate (Fig. 9d). BSR produces sulfur species
 365 reacting with iron oxyhydroxides or dissolved iron to form iron disulphides (Fig. 9e; Bonnetti et al., 2017; Machel, 2001). The incorporation of U, together with other elements (Fig. 5 and 6), may have been enhanced through their adsorption by bacterial biofilms developed at the surface of fluorite crystals (Fig. 9f). Indeed, the collomorphic fibrous texture of the external overgrowth Flog2 in the Pierre-Perthuis crystals (Fig. 3f, 4b lower right, 6 lower left and Fig. A4) could result from the development of biofilms in pore/geode linings. Although the secretion of extracellular polymeric substances leading to the
 370 formation of bacterial bio-films on pyrite has been extensively documented, their development at the surface of fluorite crystals remains largely unexplored, apart from Heim et al., (2012) who reported tubular structures within fluorite filling fractures in the Äspö diorite in Sweden, interpreted as fossilized microbial bio-films.



375 **Figure 9: Conceptual model of uranium incorporation in the fluorite lattice. a) Uranium is leached by F-rich solutions flowing through the granitic basement and reaching Fl_{geo} in the basement/cover interface, b) formation of purple layers on the fluorite surface by F-bearing fluids, c) Crystallization of Fl_{og1} , d) a local redox front is generated by BSR, e) sulfur species react with iron oxyhydroxides or dissolved Fe to form pyrite, f) uranium is incorporated in Fl_{og2} .**

380 The development of purple colorations in fluorite has long been recognized to be related to the vicinity of uranium-bearing
 minerals (Chatagnon et al., 1982; Pi et al., 2007). Bill and Calas (1978) proposed that the incorporation of Eu^{2+} , together with
 metallic calcium colloids formed by long-time irradiation, can cause purple colorations. Chatagnon et al., (1982) and later
 Kempe et al. (2002) show that such colorations have rather been caused by the reduction and stabilization of Tm and Ho in
 385 the divalent state under irradiation by the radioactive decay series of U. SR-XRF mapping focused on small well-defined purple
 growth bands (Fig. 10) shows enrichment in U and Th. Because the width of such bands is sometimes less than 10 μm , we
 have discarded the hypothesis of a coloration acquired through internal alpha irradiation (Dill and Weber, 2010; Pi et al., 2007;
 Vochten et al., 1977). Moreover, it must be noted that such coloration is not developed in the U-bearing $\text{Fl}_{\log 2}$ layer (Fig. 5 and
 6). We rather suggest that coloration was acquired through external gamma irradiation. In the following section, we will further
 discuss the spatial distribution and retention of U and Pb and the reliability of the U-Pb geochronometer in fluorite.

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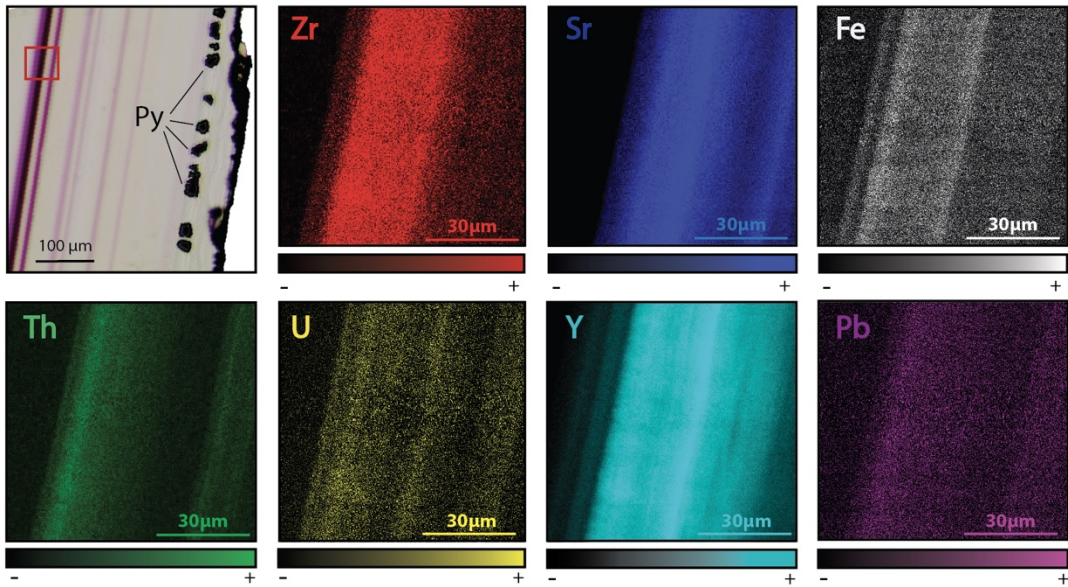


Figure 10: Transmitted plane polarized light photography from PP18S7 and corresponding Synchrotron X-ray fluorescence elemental maps. Area of XRF scan is outlined by a red box on the photography. Map dimensions are 80 x 80 μm .

395 5.2 Fluorite U-Pb geochronology

5.2.1 Retention of U in fluorite

We further discuss potential alterations of the U-Pb geochronometer by (1) the development of fractures and cleavages; (2) dissolution and re-crystallization; (3) solid-state diffusion.

(1) Fractures and cleavages

400 Fluorite is well known to develop cleavages and fractures. Although cleavages are abundant in the core of the crystals (Fl_{geo}), they are rare on the rims (Fig. 3c). Scarce fractures crosscutting Fl_{log2} were observed, causing the oxidation of pyrite (Fig. 3e). Adjacent pyrite crystals being unaffected, these fractures only generated local alterations, and were consequently avoided in LA-ICP-MS analyses (Fig. 3d).

(2) Dissolution and re-crystallization

405 Dissolution and re-precipitation of fluorite in hydrothermal deposits is common (Burisch et al., 2017). In Pierre-Perthuis, some crystals show irregular surfaces demonstrating that a late fluid event was able to slightly dissolve fluorite. These corroded crystals were not selected for LA-ICP-MS analyses. In the other crystals, the preservation of pyrite crystals, the internal fibrous texture of fluorite and the perfect geometric relationship between Fl_{geo} and Fl_{log} (Fig. 3d and 3e) demonstrate that Fl_{log2} was not dissolved and re-crystallized.

410 (3) Solid-state diffusion

Cherniak et al., (2001) determined the diffusion rates of Sr, Y, and REE in natural fluorite and concluded that no significant diffusion (< 100 µm) should occur at temperatures below 500 °C. Bosze and Rakovan, (2002), however, document diffuse boundaries between REE sectoral zoning in fluorite, interpreted as resulting from the diffusion of these elements over a distance of 200-300 µm. In Pierre-Perthuis, the preservation of ca. 10 µm thick violet bands with sharp terminations demonstrates the
415 absence of significant REE diffusion (Kempe et al., 2002). SR-XRF mapping also shows the preservation of Sr, Y, Th, Pb and U-rich bands of a thickness of ca. 10µm (Fig. 6), evidencing the absence of solid-state diffusion of these elements in our samples. Although the crystallization temperature of Fl_{log2} is unknown due to the absence of fluid inclusions, a maximum temperature of 100 °C is speculated based (1) on the fluid inclusion homogenization temperature ranging from 80 to 100°C for Fl_{geo} at Pierre-Perthuis (Gigoux et al., 2016), (2) δ³⁴S values of pyrite suggesting BSR and (3) on the fact that the
420 paleotemperatures reached by the *Assise de Chitry* Fm in the area were low during the Cenozoic period (Barbarand et al., 2013; Gigoux et al., 2016). We thus conclude that U and Pb did not diffuse within Fl_{log2}.

5.2.2 In situ LA-ICP-MS U-Pb dating

Uranium concentration in fluorite is highly variable (Piccione et al., 2019; Walter et al., 2018; Wolff et al., 2016), and is often insufficient for U-Pb dating (Piccione et al., 2019). In the Pierre-Perthuis fluorite ore, the geodic fluorite Fl_{geo}, dated at 130 ±
425 15 Ma by Sm-Nd (Gigoux et al., 2015), could not have been tested for U-Pb geochronology because it contains no U. Likewise,

Fl_{log1} is not amenable to U-Pb dating. The first fluorite generation, Fl_{leuh}, precipitated during the dissolution of the host dolomite rock, and therefore contains too many carbonate impurities that cannot be excluded from sampling in LA-ICP-MS.

430 Compared to U-Pb geochronology of other common lead-bearing minerals such as calcite (e.g., Roberts et al., 2020), fluorite U-Pb geochronology presents some additional analytical challenges. Laser induced damage in fluorite were extensively studied (De Bonis et al., 2014; Jia et al., 2005; Rafique et al., 2012; Reichling et al., 1994). Due to thermal stress and shock waves, large fractures and the ejection of solid fragments commonly occur during the first laser shots (Gogoll et al., 1996; Johansen et al., 1995). A 7s pre-ablation, excluded from data used for U-Pb age calculation, helps in getting more stable ablation rates.

435 Ablation quality depends on numerous factors including laser parameters (wavelength, fluence and repetition rate) and intrinsic crystallographic features like orientation, chemical impurities, fluid inclusions or defects (Gogoll et al., 1996). Therefore, ablation rate and quality may largely vary from one natural fluorite crystal to another.

Variations in ablation rates (crater depth/crater diameter) may generate variable downhole fractionations (DF). To minimize DF, a low laser fluence (typically 1J.cm⁻²) and a large diameter are generally preferred (Mangenot et al., 2018). However, a fluence of at least 6J.cm⁻² was required to ablate Fl_{log2}. Optical profilometry measurements of ablation pits in Fl_{log2} reveal 440 variable laser crater aspect ratios (see Fig. A1 for measured crater depths by optical profilometry). The total crater depths, including the material excluded for analysis during the 7s “pre-ablation” vary from 103 to 145 µm in Pierre-Perthuis fluorite samples and is ca. 210 µm for the HK13 fluorite (see Table A5). To correct for DF, a matrix-match primary RM can be used if laser ablation generates similar crater aspect ratios between the reference material and unknown samples (Elisha et al., 2020; Guillong et al., 2020). Such a matrix-match RM is not yet available for fluorite U-Pb geochronology (Piccione et al., 445 2019). However, even with the use of such a RM, DF correction may turn out to be incorrect because of variable crater aspect ratios in fluorite. Further research is needed to improve the precision and accuracy of fluorite U-Pb geochronology, through the systematic comparison of LA-ICP-MS crater aspect ratios in fluorite samples of known ages. In addition to DF, other sources of elemental fractionation should also be investigated, such as the influence of the chemical composition of the aerosol (Sylvester, 2008). Piccione et al. (2019) reported comparable U-Pb ages acquired on fluorite and associated nacrite, suggesting 450 that this effect may not introduce a significant bias.

5.3 U-Pb age significance and wider considerations regarding radiometric dating of ore deposits

In the absence of elemental fractionation correction against a fluorite RM, the age of the last generation of fluorite in Pierre-Perthuis ore (Fl_{log2}), 40.0 ± 1.7 Ma, may be over- or underestimated. Considering an age offset of 20% as an extreme case (Guillong et al., 2020), this age estimate still significantly differs from the Sm-Nd age of 130 ± 15 Ma (Gigoux et al., 2015) of 455 the prior geodic fluorite crystals Fl_{geo}.

Bergerat (1984) proposed an N-S compressive stress field during the Eocene through micro-structural analysis in the studied area, interpreted as the consequence of the Pyrenean compression. Evidence of late Eocene brittle deformation and associated fluid flows are also found in the eastern Paris basin, ca. 150 km north-east of Pierre-Perthuis, where calcite cementing hydraulic breccia were dated at 43 ± 1 Ma (Brigaud et al., 2020; Pagel et al., 2018). Volcanic activity in the northern part of the French Massif Central has been reported by Bellon et al., (1974) and Lefort and Agarwal, (2002) during this period, at about 80-120km south-east of the study area. In conclusion, a local compressive stress field related to the propagation of N-S Pyrenean compression in the intraplate domain may have generated fluid flows and the crystallization of Fl_{log2} .

In Pierre-Perthuis, the geodic fluorite crystals Fl_{geo} yield a Sm-Nd age of 130 ± 15 Ma (Gigoux et al., 2015). Here, we show that later fluid flows took place during the Cenozoic period, recorded as rims overlaying Fl_{geo} . Hence, at least two mineralizing events occur during a time period of ca. 90 Ma.

In some ore deposits, fluorite Sm-Nd dating is the only available geochronometer. However, errorochrons and imprecise ages may occur. In such cases, U-Pb geochronology of authigenic phases with elevated common Pb such as calcite (Rasbury and Cole, 2009), hematite (Walter et al., 2018), chalcedony (Nuriel et al., 2011) or nacrite (Piccione et al., 2019) offers new perspectives to gain knowledge on the successive chronology of mineralizing fluid flows. In Pierre-Perthuis however, as in some other F-Ba deposits, these minerals are absent. A late Eocene age estimate for the last fluorite-mineralizing event reveals that the ore-formation is polygenic. Fluorite was deposited during several successive fluid flows induced by the reactivation of faults affecting the granitic basement. Small-scale radiometric dating of single growth bands in authigenic phases makes it possible to determine the chronology of these multiple crystallization events. Future research should be conducted to test the U-Pb geochronometer in fluorite crystals dated by Sm-Nd or other direct ((U–Th–Sm)/He) or indirect methods.

6 Conclusion

We have explored the U-Pb geochronometer in fluorite. In the Pierre-Perthuis F-Ba deposit, uranium was found in the most external rim of fluorite crystals. SR-XRF mapping demonstrates the preservation and retention of micrometer-thick zonation in uranium concentration in fluorite. Petrographic observations coupled to induced fission tracks and SR-XRF mapping show that uranium is incorporated in the fluorite crystal lattice by elemental substitution. The incorporation of uranium is related to bacterial sulphate reduction, evidenced by globular pyrite inclusions. Relying on careful and detailed petrography, we show that such growth bands can be dated by U-Pb geochronology. The Tera-Wasserburg lower intercept yields an age of 40.0 ± 1.7 Ma, not corrected for matrix-related elemental fractionation. This age estimate corresponds to the last minor fluorite-mineralizing event. Since the crystal cores were previously dated at 130 ± 15 Ma by Sm-Nd, at least two episodic fluid flows occurred during ca. 90 Ma. Future research should be conducted to correct for $^{238}\text{U}/^{206}\text{Pb}$ fractionation during LA-ICP-MS analysis in fluorite through the definition of a fluorite reference material.

Appendix A

Uranium concentration measured by counted fission tracks

The range of latent track lengths of fission particles depending on the density of materials, the correction factor between two matrixes is given by Enkelmann et al., 2005. The latent fission-track lengths corresponding to the combined ranges of the light (Lt) and heavy (Hv) nuclide ejected in opposite directions from the fission of ²³⁵U after capture of a thermal neutron (Jonckheere, 2003) have been simulated by SRIM® software (Ziegler et al., 2010). The numerical values for mass and energy of these nuclides are taken from Kahn et Forgue (1967), glass standard reference CN5 density from De Corte et al., 1998 and all data and simulations results are summarized in Table A1.

(a) Projectiles	Z	M _A (amu)	E ₀ (MeV)
Hv (Xe)	54	138.1	69.6
Lt (Sr)	38	95.34	100.1
(b) Matrices	Formula	d (g.cm3)	
Glass CN5	SiO ₂	2.45	
Fluorite	CaF ₂	3.18	
(c) Simulations	Range Hv (μm)	Range Lt (μm)	[2R] (μm)
Glass CN5	10.5	14.7	25.2
Fluorite	9.5	12.9	22.4
(d) Correction factor	d _{Glass CN5} /d _{Fluorite}	[2R] _{Glass CN5} / [2R] _{Fluorite})	F _{corr}
Glass CN5/Fluorite	0.770	1.125	0.866

Table A1: Data used to calculate the correction factor for uranium concentration in fluorite using induced fission track mapping in SRIM® software. a) mass (M_A) and energy (E₀) of heavy (Hv) and light (Lt) nuclides produced by the fission of ²³⁵U, b) Formula and density (d) of the two simulated materials, c) Latent track lengths (2R) obtained by adding both nuclides simulated range lengths, (d) uranium concentration correction factor (F_{corr}) calculated for fluorite with glass standard material (Enkelmann et al., 2005).

Sample	zone	counted fission tracks	area (μm^2)	U concentration (ppm)	U concentration corrected (F_{corr})	U average concentration /sample (ppm)
CORNING CN5	-	131	10^4	12.17	-	-
PP1802-1	1	57	3000	18	16	16
	2	57		18	16	
	3	54		17	15	
	4	63		20	17	
	5	62		20	17	
	6	58		19	16	
	7	52		17	14	
	8	60		19	17	
PP1802-2	1	67	2500	26	23	25
	2	70		27	24	
	3	76		30	26	
	4	85		33	29	
	5	76		30	26	
PP1802-4	1	56	1500	38	33	26
	1 (rim)	22		15	13	
	2	58		39	34	
	2 (rim)	31		21	18	
	3	47		32	28	
	3 (rim)	25		17	15	
	4	86		58	50	
	4 (rim)	22		15	13	
	5	63		43	37	
	5 (rim)	29		20	17	
PP1801-2	1	36	3000	13	11	7
	2	28		10	8	
	3	26		9	8	
	4	13		5	4	
	5	22		8	7	
	6	25		9	8	
	7	33		11	10	
	8	13		5	4	
CORNING CN5	-	94	10^4	12.17	-	-

Table A2: Fission tracks analytical results. Counted areas are distributed along $F_{\log 2}$ on different samples. The correcting factor employed for U concentration calculation on each sample depends on the counted area and followed a bracketing method (thanks to the two CN5 standard certified at 12.17ppm).

Estimation of the penetration depth of SR-XRF

Element	X-ray line, (in keV)	CaF ₂ , information depth, (in μm)
S	2,3	5,5
Ca	3,691	20
Fe	6,403	18
Y	14,957	189
Sr	14,164	162
Zr	15,774	220
U	13,613	145
Pb	10,549	70
Th	12,966	126

Table A3: Information depth estimation (in μm) and measured X-ray line (in keV) of the SR-XRF method in fluorite for each element.

LA-ICP-MS U-Pb dating parameters

Laboratory and Sample Preparation	
Laboratory name	Géosciences Paris Sud (GEOPS). Université Paris-Saclay. Orsay. France
Sample type/mineral	Fluorite
Sample preparation	Fluorite crystals mounted in epoxy resin
Imaging	Optical microscopy using transmitted light
Laser ablation system	
Make. Model and type	193nm ArF (Teledyne Photon Machines)
Ablation cell	HelEx
Laser wavelength (nm)	193 m
Pulse width (ns)	5 ns
Fluence (J.cm-2)	6.25 J.cm ⁻²
Repetition rate (Hz)	10 Hz
Pre-ablation	each spot during 7 s
Ablation duration (secs)	30 s
Spot size (mm)	150 μm
Sampling mode / pattern	Static spot ablation
Carrier gas	He
Cell carrier gas flow (l.min ⁻¹)	Helium Lage volume: 0.5 l.min ⁻¹ Inner cup: 0.375 l.min ⁻¹
ICPMS instrument	

ICPMS instrument Make. Model & type	ThermoScientific Element XR
Sample introduction	Ablation aerosol
RF power (W)	1175 W
Make-up gas flow in ablation funnel (l.min ⁻¹)	Ar=0.950 to 1 l.min ⁻¹
Detection system	Ion counter
Masses measured	206. 207. 208. 232. 238
Average gas background (cps) 2019-12-19	12 for 206. 10 for 207. 26 for 208. 0.1 for 232. 0 for 238
Average gas background (cps) 2019-12-20	20 for 206. 17 for 207. 53 for 208. 0.1 for 232. 0 for 238
Integration time per peak (ms)	10 ms for ²⁰⁸ Pb, ²³² Th, ²³⁸ U. of 30 ms for ²⁰⁶ Pb and of 40 ms for ²⁰⁷ Pb by 70 runs
Total integration time per reading (secs)	0.1 s
IC Dead time (ns)	30 ns
Signal strength at ICPMS tuned conditions	Th/U=1.02 248ThO/ ²³² Th below 0.3%
Data Processing	
Data acquisition	Fully automated mode overnight in sequences of 399 analysis maximum
Gas blank	30 s background. 30 s sample ablation and 30 s washout
Calibration strategy	NIST614 for Pb-Pb. calcite WC-1 for U-Pb. secondary reference materials: calcite Duff Brown Tank (DBT). calcite breccia AUG-B6 and fluorite HK13
Reference Material info	WC-1 age: 254.4±6.4 Ma (Roberts et al., 2017). Duff Brown Tank age: 64±0.7 Ma (Hill et al., 2016). Calcite breccia AUG-B6 age: 43±1 Ma (Pagel et al., 2018) and fluorite HK13 age: 290±10 Ma (Wolff et al., 2016)
Data processing package used / Correction for LIEF	Iolite to calculate uncertainties. no down-hole fractionation correction
Mass discrimination	²⁰⁷ Pb/ ²⁰⁶ Pb normalization to NIST614. ²⁰⁶ Pb/ ²³⁸ U normalization to WC-1
Common-Pb correction. composition and uncertainty	No common-Pb correction applied
Uncertainty level & propagation	Ages in the data table are quoted at 2sigma (2σ) absolute. uncertainty propagation by quadratic addition
Quality control / Validation 2019-12-19	Measurements of WC-1 age = 167.04 ± 3.7 Ma. Duff Brown Tank (DBT) age = 63.75 ± 2.2 Ma. AUG-B6 age = 40.7± 1.6 Ma and NIST614 were done along with samples throughout the analytical session
Quality control / Validation 2019-12-20	Measurements of WC-1 age = 148.02 ± 2.3 Ma. Duff Brown Tank (DBT) age = 64.57 ± 2.9 Ma. AUG-B6 age = 44.45 ± 2.0 Ma. “HK13” fluorite was dated at 285.9 ± 29.3 [30.9] Ma and NIST614 were done along with samples throughout the analytical session

Table A4: Data for fluorite LA-ICP-MS U-Pb analysis at University Paris-Saclay, GEOPS laboratory.

Depth and diameter of laser craters

- 515 3D images of laser craters have been acquired with an optical profilometer 3D Contour GT-X (Bruker Corp., Billerica, MA, USA) in VSI (Vertical Shift Interference) mode at GeePs laboratory, Université Paris Saclay. This instrument produces 3D images of a surface and also depth profiles on chosen axes thanks to Vision® analysis software (VISIONx Inc., Pointe Claire, Quebec, Canada). The VSI mode is based on white light vertical scanning interferometry, with measurable topography up to

1 mm and a vertical resolution of 3 nm. Lateral resolution measurements are a function of the objective magnification (a $\times 50$ magnification was used for this study) giving a lateral resolution of $0.2\mu\text{m}$.

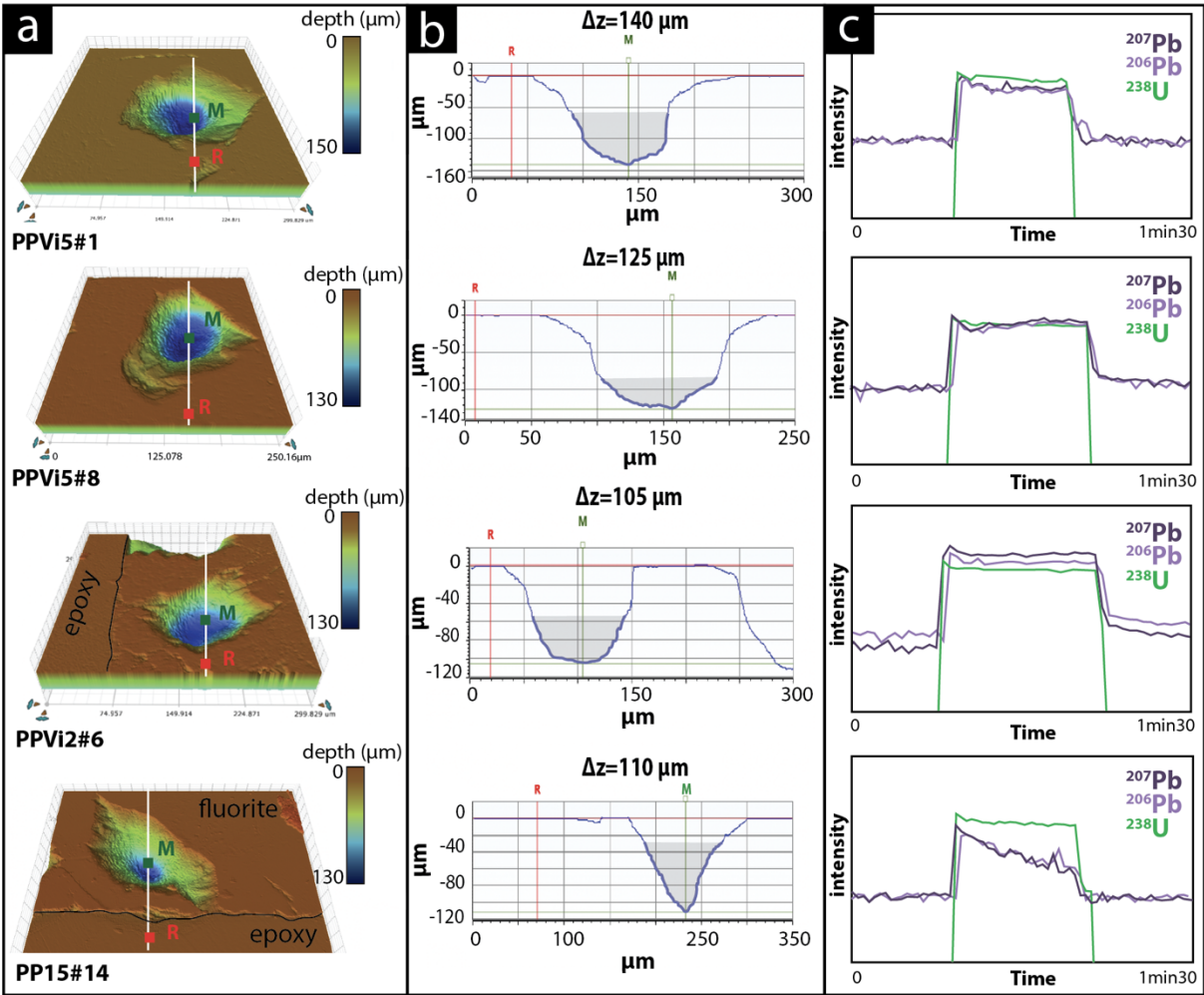


Figure A1: Various crater aspects induced by laser ablation on different fluorite sample a) 3D surface images of craters reconstructed by optical profilometer, b) corresponding 2D axial section with the measurement of crater depth thanks to two markers: R (the average value of the planar surface around the crater) and M (the deepest point in the crater) expressed as Δz in μm , in grey is illustrated the volume below a crater diameter of $85\mu\text{m}$, c) the corresponding LA-ICP-MS signal intensity of ^{238}U , ^{206}Pb and ^{207}Pb during the laser ablation. Analyses PPVi5#8, PPVi5#1 and PPVi2#6 were included in the Tera-Wasserburg diagrams, while PP15#14 is an example of analysis discarded due to variable $^{238}\text{U}/^{206}\text{Pb}$.

Sample	Total depth, including pre-ablation (μm)	volume (μm^3)
PPVi2_#13	130	574842
PPVi2_#19	116	675090

530

PPVi2_#24	145	818605
PPVi2_#25	105	626830
average	124	673842
PPVi5_#1	140	875287
PPVi5_#2	140	912850
PPVi5_#3	120	834602
PPVi5_#8	125	757431
average	131	845043
PP1802_#6	103	524342
PP1802_#9	110	507250
PP1802_#11	122	549564
PP1802_#12	132	621271
average	117	550607
PP15_#3	105	249329
PP15_#9	110	329296
PP15_#14	110	360308
PP15_#15	130	451288
average	114	347555
HK13#1#5	210	N.D.

Table A5: Statistical analysis of crater depths induced by laser ablation for U-Pb dating by optical profilometer. The total crater depths and volumes include the 7s pre-ablation.

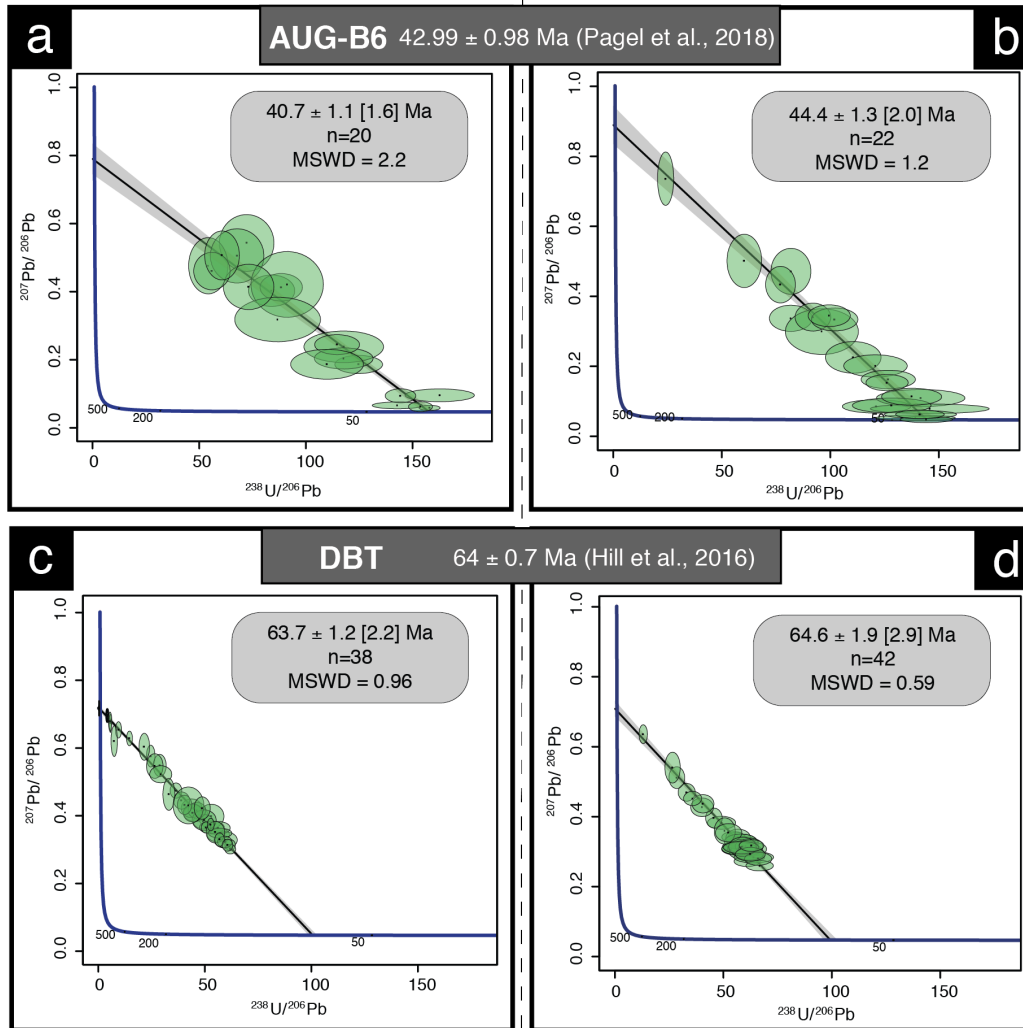


Figure A2: Tera-Wasserburg diagrams of secondary calcite reference materials. (a,b) Tera-Wasserburg diagrams displaying corrected ages for the calcite of the Gondrecourt graben AUG-B6 measured during the two analytical sessions, (c,d) Tera-Wasserburg diagram displaying the Duff Brown Tank corrected ages measured during the two analytical sessions

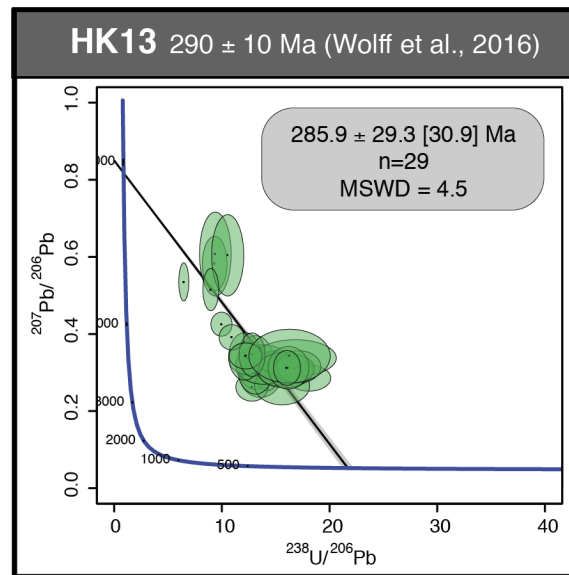
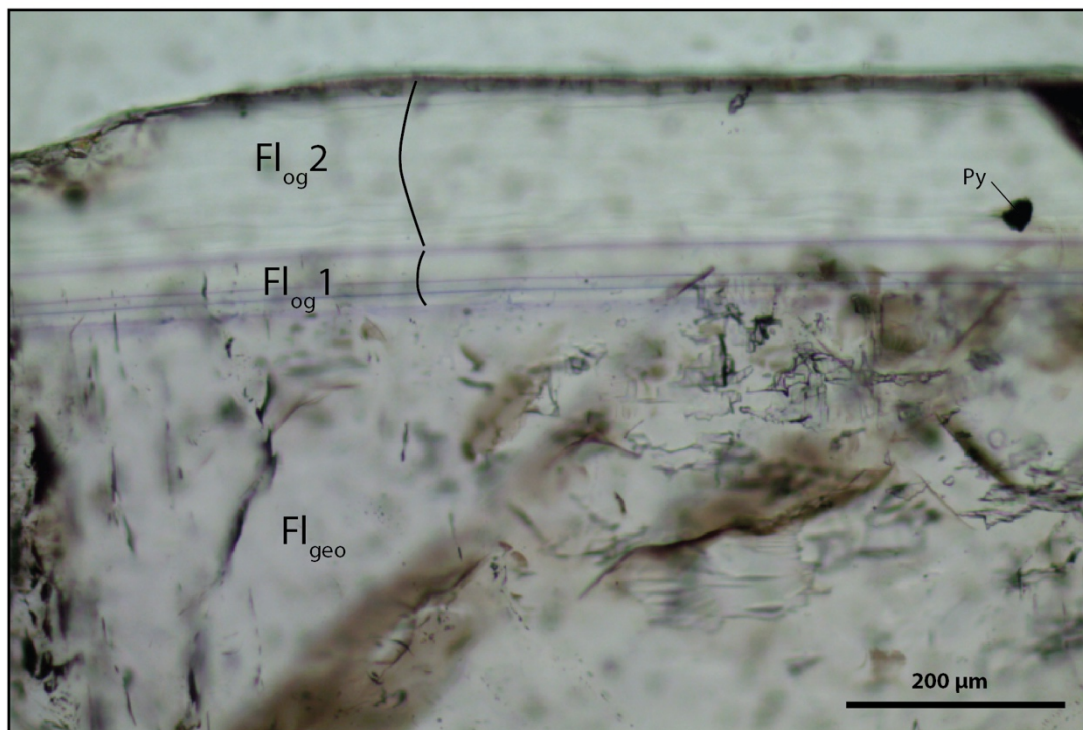


Figure A3: Tera-Wasserburg diagrams of secondary fluorite reference materials, “HK13” fluorite dated at 290 ± 10 Ma by Wolff et al., 2016.

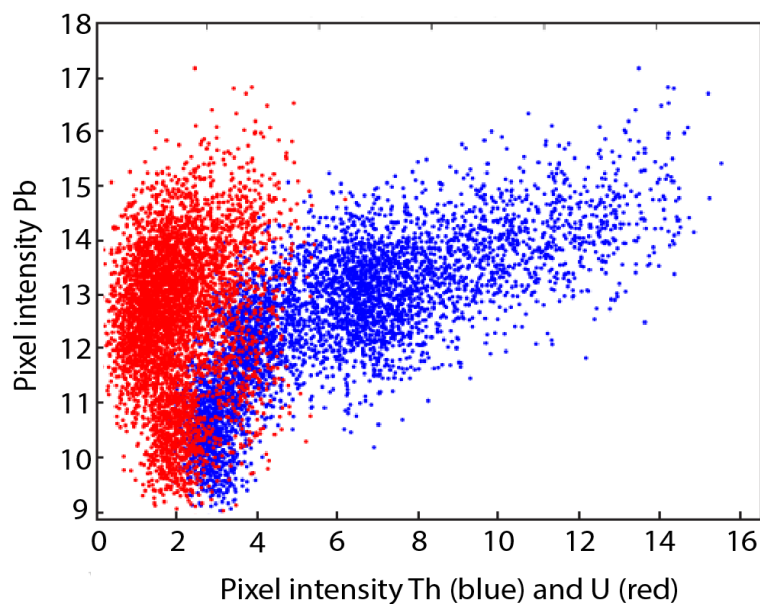
Petrographic observations



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Figure A4: Microphotography of the fluorite stages from the Pierre-Perthuis ore showing the contrast in fluid inclusions concentration between Fl_{geo} and Fl_{og}. Py: Pyrite.

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550 **Figure A5: Scatter plot of lead as function of thorium (blue) and uranium (red) in pixel intensities, corresponding to the scanned area illustrated in Fig. 6.**

Author contribution

Julius Nouet and Claire Boukari were involved in the preparation of the fluorite samples for SEM observations, SR-XRF and induced fission track mapping. Andr a Somogyi directed the elemental mapping experiments on the Nanoscopium beamline at synchrotron SOLEIL, and supervised SR-XRF data interpretation. Aurore Br zard-Oudot carried out the measurements by 3D optical profilometry at Group of electrical engineering Paris (GeePs) laboratory. Jocelyn Barbarand, Benjamin Brigaud and Maurice Pagel were involved in field work, data interpretation and contributed to the improvement of the manuscript. Louise Lenoir and Thomas Blaise participated in every step of the study and prepared the manuscript.

Competing interests

560 The authors declare that they have no conflict of interest.

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