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Impact of incipient weathering on uranium migration in granitic waste rock piles from former U mines (Limousin, France)



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ABSTRACT

This study investigates the impact of incipient weathering on uranium (U) migration at the surface of granitic waste rock piles from former U mines in Limousin (France) operated half a century ago. We used a multi-scale mineralogical and geochemical approach to evaluate the impact of incipient weathering on uranium mobility in a chaotic waste rock pile. The surficial part of five former mines located in the "La Crouzille" and "La Creuse" mining districts was investigated. Four groups of samples were defined based on field observations and mineralogy, granites, lamprophyres, hydrothermally altered rocks and weathering products. Petrology, mineralogy and geochemistry investigations show the importance of mineralogical and geochemical transformations during incipient weathering over a few decades. The Chemical Index of Alteration (CIA) is successfully used to assess the weathering state of samples in the waste rock pile chaotic deposit. The mean CIA value is around 60 for unweathered/incipiently weathered granites whereas highly weathered samples have CIA around 70. This means that no significant major element loss occurs in the granitic rocks suffering incipient weathering. Various micron-size secondary U minerals consist in a large diversity of oxidized uranyl minerals (uranophane, uranyl sulfate, curite, autunite and other uranyl phosphates), with dominant uranyl phosphates, indicating local U mobility in the waste rock pile. Autunites micromorphology indicates different degrees of weathering of the samples in which it crystallizes. U is primarily concentrated in the clay fraction (< $2 \mu m$), associated with secondary minerals from granite alteration (clay minerals such as smectite and vermiculite, Fe-oxides and phosphate minerals) and, to a limited extent, relict primary minerals. These weathering products contribute to U (VI) stabilization under oxidizing conditions.

1. Introduction

Post-mining environmental monitoring has become an important part of the mining-life cycle (Jamieson et al., 2015). In Western Europe, granitic intrusions were sites of intense mining activity in the 20th Century. In France, out of nearly 250 former U mining sites, 50 are located in granitic massifs of the Limousin region, which hosts 57 Mt of waste rocks (AREVA, 2004). With a high specific surface, granitic waste rocks suffer enhanced weathering rates compared to regular outcropping granites. Recent interest has arisen concerning the environmental impact of granites, among which the sorption of uranium on granite for modeling high-level radioactive waste geological repositories (Lee et al., 2011; Jin et al., 2016). These works showed that, in the pH range 4–9, U(VI) sorption increases up to 90% within 72 h. Investigation of the environmental impact of uranium released by the industrial processing of granitic materials (scraps, water and slurries) also shows modification of U speciation with increasing pH (Guillén et al., 2014). In addition, granite alteration studies by column acid leaching have demonstrated that weathering affects U distribution (Guthrie and Kleeman, 1986). A broad range of fields of environmental importance is concerned by granite weathering. Understanding the processes governing U migration under these conditions is thus important to model source terms for a potential U environmental impact.

Granite weathering has been extensively studied in natural regoliths (Gilkes et al., 1973; Nesbitt and Young, 1989; White and Brantley, 1995; Braga et al., 2002; Takaya, 2014; Scarciglia et al., 2016) and in monitored waste rock piles (Smith et al., 2013). The weathering rate of granite primarily depends on the weathering rate of its least stable minerals, biotite and plagioclase (Gilkes et al., 1973; White and Brantley, 1995; Takaya, 2014) and incipient weathering is mainly

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controlled by sulfide oxidation and Al and Fe-oxyhydroxide precipitation (Smith et al., 2013). The weathering intensity is quantified by several weathering indexes (Price and Velbel, 2003). Among these, the "Chemical Index of Alteration" (CIA), defined by the ratio $Al_2O_3 /$ ($Al_2O_3 + CaO + Na_2O + K_2O$) (Nesbitt and Young, 1989; Scarciglia et al., 2016) is frequently used to assess the weathering state of basaltic or granitic regoliths. It is a good proxy of continental crust weathering because it focuses on feldspar alteration, one of the most common minerals of the continental crust.

Field based studies show the liberation of potential contaminants of concern (including U) from waste rocks from former U mines during exposure to weathering under oxidizing conditions after a few decades (Tran et al., 2003; Kipp et al., 2009; Parbhakar-Fox et al., 2014; Cabral Pinto et al., 2014; Neiva et al., 2016). These studies highlight the local mobility of U(VI) and its sorption on nano/micro-divided weathering minerals, such as clay minerals or Fe-oxides, or U(VI) sequestration by phosphate minerals. In natural regolith profiles, similar trends are observed: trace elements, including U(VI), are found along with phosphate minerals, clay minerals and Fe-oxides (Jerden et al., 2003; Taboada et al., 2006; Regenspurg et al., 2010). Despite the diversity of U(VI) minerals, when phosphate is present in natural regoliths, U bearing phosphate minerals are predominant (Jerden et al., 2003; Jerden and Sinha, 2003; Jerden and Sinha, 2006; Knox et al., 2006). U mobility is determined using various multi-scale approaches: from the field scale, identifying faulted structures responsible for U re-concentration after hydrothermal episodes (Lespinasse and Cathelineau, 1990) to the molecular scale, focusing on U speciation (Morris et al., 1996; Allard et al., 1999; Kelly, 2010; Morin et al., 2016).

A preliminary study (Boekhout et al., 2015) was performed on a trench dug in a granitic waste rock pile in the former Vieilles Sagnes uranium mine (Limousin, France), to evaluate the feasibility of our approach. The present study focuses on surficial incipient weathering at the very surface of five granitic waste rock piles of two former major U mining districts; "La Crouzille" and "La Creuse", Limousin, France (International Atomic Energy Agency, 2009). This region is one of the largest vein-type uranium districts in Europe, and produced 23,300 t U at an average grade of 0.2%, between 1948 and 1995, resulting in 57 Mt of granitic waste rocks that were left on site and partly re-vegetated (IRSN, 2007). Previous environmental studies in nearby areas have demonstrated uranium mobility and its accumulation in certain levels of peat soils (up to 12,000 ppm) by precipitation of colloids of Fe, Al and OM or P-Al-Fe-Si (Wang et al., 2013; Leermakers et al., 2016). In the alpine region (Europe), natural soils arising from granite concentrate up to 4000 ppm of U, despite the absence of mining activity in the area (Regenspurg et al., 2010). Weathering products of granite limit U migration by sorption on Fe-oxyhydroxides (Leermakers et al., 2016) and clay minerals. The formation of uranyl phosphates is also observed (Boekhout et al., 2015). These phosphates are an efficient trap for uranium sequestration (Munasinghe et al., 2015).

The present study investigates U migration in the surficial part (5 to 30 cm deep) of 5 granitic waste rock piles from former U mines in the Limousin region, France, with an intense mining activity, 24 to 66 years ago. The granitic boulders investigated (10 cm to 30 cm large) present, from the rim to the core, incipient weathering features and show the influence of the previous geological history and the vegetation development. CIA parameters show coherent weathering trends from the granitic blocks to the underlying technosoil. Incipient weathering is related with CIA values between 55 and 65, whereas highly weathered samples have CIA around 70. Mechanical and chemical alteration of waste rock piles affects the soils and peat soils downslope. We discuss the mobilization of uranium, the first trapping processes associated to mineralogical and geochemical transformations and the preferential occurrence of specific secondary U minerals, such as uranyl phosphates.

2. Materials and methods

2.1. Site description and sampling

Fifty years of mining activity in vein-type granite-hosted U deposits in Limousin resulted in 15 to 30 m high waste rock piles of dominantly granitic boulders (10 cm to 1 m large), subject to weathering and partly re-vegetated (with ferns and 2-5 m high birch trees). We sampled the waste rock piles from 4 selected former mining sites in the "La Crouzille" former mining district (Limousin, France), located in the European Variscan belt, in a two-mica granite massif (International Atomic Energy Agency, 2009). A brief description of the regional geological background is given in Appendix A. Additionally, the waste rock pile of the Hyverneresse mine, in the "La Creuse" mining district, Limousin, France (located in the Millevaches granite, a distinct, though contemporary, two-mica granite massif) was sampled. Hyverneresse is located in a felsic facies, principally composed of leucogranite (Stussi and Cuney, 1993). Granite mineralogy and petrology are similar in both the "La Crouzille" and "La Creuse", mining districts (Leroy, 1984; Bonhomme et al., 1987). In both districts, the ore is mainly composed of uraninite, pitchblende and coffinite.

Most waste rocks correspond to non-mineralized granites from overburden, extracted to access the ore veins. The average U concentration in these granites is 20 ppm (Cuney, 2014; Ielsch et al., 2016), in agreement with the regional U background of Limousin granites. These two-mica granites contain around 36% quartz, 27% albite, 27% orthoclase and 10% biotite and muscovite (Dahlkamp, 1993). Hydrothermal episodes or supergene alteration induce neoformation of calcite and pyrite (Leroy, 1978), jarosite and clay minerals (McLemore et al., 2006; Salama et al., 2016). A minor part of the waste rocks corresponds to low-grade ore below the "cut-off grade", with a bulk U concentration, from 100 to 300 ppm (IRSN, 2007). This cut-off grade was assessed on a large volume, 0.4 m^3 (AREVA, 2004), whereas hand sampling concerns volumes 10^3 times smaller, 0.4 dm^3 . Therefore isolated samples may exceed the average cut-off value of 300 ppm.

Five post-mining sites were studied: Bellezane, Fanay, Hyverneresse, Margnac and Pény (Appendix B). Sampling was restricted to the surficial 5 to 30 cm of the waste rock piles. Piles can reach 20 m high, with a slope of 10 to 1000% (Table S1 in the Supplementary material). The Pény and Hyverneresse slopes are steep (over 500%) and vegetation is scarce (Fig. 1a, c). The Fanay slope (75%) is more vegetated (Fig. 1b). On the contrary the Bellezane slope (75%) has no vegetation. Bellezane site is a former open pit mine, with walls showing lamprophyre intrusions (Fig. 1d). Lamprophyre and granite have similar chemical composition. Due to their micro-granular texture, lamprophyres are more sensitive to weathering than granites (Rock, 1984). In this paper surficial lamprophyre represents a reference of intense supergene alteration.

Sampling focused on samples with a high radiometric signal in order to get the highest U concentration possible. The radiometric signal (dose rate) ranges from $0.3 \,\mu \text{Sv} \cdot \text{h}^{-1}$ to $2.4 \,\mu \text{Sv} \cdot \text{h}^{-1}$. Fifty samples representative of the diversity in mineralogy, morphology, and weathering degrees were selected and sorted in four groups: (Table S2 in the Supplementary material):

- Granites with apparent weak alteration (27 samples): samples are decimetric, hard blocks of granites with visible alteration features. Some have cracks and fractures, as well as color changes from fresh pink in the bulk to orange and brown at the boulder surface due to oxidation. Most blocks composing the waste rock pile belong to this category (Fig. 1a, b and c).
- Lamprophyres (4 samples): at the open pit mine of Bellezane, two different outcrops of lamprophyre dykes have been sampled at different levels of the wall of the mine (not from the pile). The deepest outcrop (10 m below surface) was partially affected by



Fig. 1. Pictures showing the variety of samples collected. (a) Vegetalized granitic rock heap at Pény. (b) Well developed vegetation on Fanay's upper waste rock pile. (c) Ferrihydrite pools (Fh p.) downslope Hyverneresse waste rock pile. The waste rock pile (indicated by yellow dotted lines) shows vegetation development. (d) Lower lamprophyre dike at Bellezane. The dyke is indicated in the granitic massif by red dotted lines. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

hydrothermal alteration (Fig. 1d). The surficial part of the dyke was affected by intense supergene alteration. Lamprophyre blocks are scarce in the waste rock pile and generally highly weathered. Therefore they were sampled in the wall-rock for a representative sampling.

- Hydrothermally altered rocks (8 samples) are divided in two subgroups:
 - o a sub-group of friable coarse grained granite containing calcite (3 samples).
 - o a sub-group of hard fine grained rocks containing sulfides (5 samples) with a yellow coloration.
- Weathering products (15 samples) can be divided in 4 sub-groups: o Arenization matrix of the weathered rim of the blocks is found as silty-clay weathering aureoles of submetric granitic blocks. Weathering associated with fracturing has led to the arenization of the waste rocks.
 - o Protosoils correspond to the incipient formation of centimetric sandy-silty horizon at the surface of the waste rock pile associated with moss growth and located below birch trees mid slope of the pile.
 - o Technosoils are located on flat surfaces either at the top or at the bottom of the piles. They consist of a 10 to 20 cm thick siltyclayey brown soil with a variable humic fraction. They are eventually mixed with soil cover from the overburden, after site

rehabilitation.

o Ferrihydrite-rich orange and muddy meter-wide pools form downslope. These pools, distinct from the rills, indicate a Fe migration and they are specific to the waste rock pile of Hyverneresse (Fig. 1c).

2.2. Analytical procedures

XRD analyses were performed using an X'Pert Pro Panalytical diffractometer with Ni filtered CuK α radiation at 40 kV and 40 mA. All samples were ground with an agate mortar and pestle prior to bulk analyses. The < 2 µm fraction was obtained by size-fractionation of bulk samples in deionized water according to Stoke's law. Disoriented powders of all samples were scanned from 3° to 75° (20) with a counting time of 520 s every 0.0167°, in an aluminum rotating sampleholder. For clay mineral identification, the < 2 µm fractions were deposited by evaporation from deionized water on glass–disk holders. Oriented air-dried, glycolated and heated (550 °C for 3 h) samples were scanned from 3° to 15° with a counting time of 270 s every 0.0167°.

Dry granulometric separations were performed with a set of sieves (1 cm, 2 mm, 63 μ m) on 7 samples (Table S5). Wet granulometric separations of the "clay fraction" (< 2 μ m) were performed on 9 bulk samples (Tables S4). 10 g of bulk sample was mixed with water and ultrasound for 10 min in order to separate agglomerated minerals. After 5 h sedimentation, in a total of 1000 ml of water, the supernatant was siphoned out and centrifuged at 4000 rpm for 60 min. The dried sample is referred to as the "clay fraction" (< 2 μ m), following the international nomenclature.

Thin sections were manufactured from the core of blocks 10 to 20 cm large. They were made at the Institut de Recherche pour le Développement (IRD), Bondy research center and studied with an optical microscope Olympus, BH-2.

Scanning Electron Microscopy (SEM) analyses were conducted with a Zeiss ULTRA55 with an acceleration voltage of 15 keV and a working distance of 7.5 mm. Micromorphological observations were made with secondary and backscattered electrons detectors, and an Energy Dispersive Spectroscopy (EDS) microprobe.

Chemical analyses were obtained by Inductively Coupled Plasma-Mass Spectrometry (ICP-MS) on an Agilent 7500 CX[®] at IRD Bondy Research center, for major elements (detection limits in ppm between brackets): Na(0.05), Mg(0.05), Al(0.8), P (0.02), K(0.2), Ca(0.45), Ti (0.05), Fe(0.9), Mn(0.005); and trace elements (detection limits in ppb between brackets): Ba(2.5), Ce(1.0), La(0.5), Pb(0.2), Th(0.5), U(2.5) and Zr(0.9). ICP-MS measurements were performed after multi-acid digestions (HNO₃–HF–H₃BO₃). The Si volatile component is lost during the evaporation. International reference materials (GSN and MAN) were regularly measured for quantitative accuracy estimation. Values given are issued from three replicates to insure accuracy and precision of the data. For all trace elements the precision was below 5% (RSD).

Sulfur measurements were conducted by pyro-hydrolysis at the Institut de Physique du Globe de Paris (IPGP). Values are given \pm 0.01%. Loss on Ignition (L.O.I.) measurements were carried out according to Lechler and Desilets (1987) procedure for igneous rocks. Values are given \pm 0.1%. The amount analyzed was extracted from 100 g of sample previously ground in order to get representative values.

3. Results

3.1. Incipient weathering identification using mineralogical and geochemical tools

3.1.1. Waste rock mineralogy

Micromorphology observations on thin sections (Fig. 2) and XRD mineralogy (Fig. S1 in the supplementary material) help determine the origin of the alteration (weathering, or hydrothermalism) within the four groups of samples (Table S2 in Supplementary material):

3.1.1.1. Granites

The granitic group can be subdivided in two groups, depending on the alteration state: "no or low clay minerals content" and "significant clay minerals content". The clay minerals identified are chlorite (Fig. S1-Ia in the Supplementary material) and/or kaolinite. Granitic samples show cemented mineral grains (Fig. 2a): quartz, mostly unweathered feldspars, partially weathered biotite, fresh muscovite and accessory minerals (zircons and monazites). Chlorite alteromorph after biotite (i.e. partial transformation of biotite minerals to chlorite, with conservation of the initial biotite's shape) indicates incipient weathering (Fig. 2b). Radiation aureoles evidence accessory radioactive minerals included in biotites such as zircons and monazites (Fig. 2b). Incipient weathering affects micas and feldspars with the formation of micrometric Fe-oxides veins (Fig. 2c).

3.1.1.2. Lamprophyres (Fig. S1-II in the Supplementary material). The lower part of the lamprophyre dyke mineral assemblage (Fig. 1d) is composed of plagioclase, K-feldspar, quartz, smectite and chlorite with scarce micas and amphiboles (Fig. 2f and Fig. S1-II-d in the Supplementary material). The matrix consists of clay minerals and Feoxides (Fig. 2f). In the surficial part of the same dyke, the occurrence of vermiculite, interstratified clay minerals and rare relicts of plagioclase indicates an intensely weathered state (Fig. S1-II-e f and g in the Supplementary material), observed in the most powdery part of the dyke.

3.1.1.3. Hydrothermally altered rocks

The calcite-rich rocks (Leroy, 1978) are composed of calcite, smectite, rare biotite and muscovite (Fig. S1-III in the Supplementary material). The most friable sample is the most altered, kaolinite occurrence suggests recent weathering. Sulfide-rich rocks contain quartz similar to an episyenite facies. Large areas ($600 \ \mu m^2$) of clay minerals and Fe-oxides occur in the matrix (Fig. 2d). They are affected by weathering down to the core of 20 cm blocks, as shown by clay mineral alteromorph after feldspar (Fig. 2e), pyrite dissolution figures (Fig. 3c) and jarosite neoformation (Fig. 3a) suggesting pyrite oxidation.

3.1.1.4. Weathering products (Fig. S1-1 in the Supplementary material). The mineral paragenesis of the granitic arena includes smectite, chlorite, kaolinite and interstratified clay minerals (FIG. S1-Ib in the Supplementary material) with relict primary minerals. Protosoil samples contain quartz, K-feldspars and plagioclases, micas, Fe-oxides, smectite, kaolinite (Fig. S1-Ic in the Supplementary material) and jarosite. Technosoils have a similar mineralogical signature, with a variable content of interstratified clay minerals and no or rare jarosite. They present a high content of organic matter, with an extreme example represented by a humic horizon, occurring at the bottom of the Fanay pile (Table S3 in the Supplementary material). The ferrihydrite encountered in the pools is a 2-line ferrihydrite, with two broad bands corresponding to d-spacings of 1.5 and 2.5 Å.

Fig. 2. Pictures under optical microscope. Abbreviations list: Bt = biotite, Chl = chlorite, Vrm = vermiculite Cl = clay minerals, FeOx = iron oxide, Fsp = feldspar, M = mica, Py = pyrite, Qz = quartz, PPL: Plane polarized light, XPL: Cross polarized light.

a) Lesser altered granite from Margnac, XPL. Primary minerals: Qz, Fsp and M. b) Chlorite alteromorph after biotite in a granite from Bellezane, PPL. Biotite shows radiation aureoles. c) Close up of (a), PPL: iron oxide veins in mica. d) Clay and iron oxide rich matrix in a granite from the sulfide-rich group from Fanay, XPL. e) Pyrite and iron oxides in a granite from the sulfide-rich group from Fanay, PPL. f) Lamprophyre sample, from the lower part of the dyke, showing a clay and quartz rich matrix PPL.





Fig. 3. SEM pictures of rock chips: (a) Backscattered electron image. Iron and potassium sulfate (Jarosite: Jrs) rosette formation on mica (M) and feldspar background in a sulfiderich sample from Fanay. (b) Secondary electron image of lichen growth at the surface of micas. A monazite (Mnz) crystal is visible. (c) Secondary electron image of pyrites (Py) in smectite (Sme) and mica background in a sulfide-rich sample from Fanay.

Mineralogy along with micromorphology observations helps determine the weathering impact on the waste rock piles. In the coarse groundmass of the arena, dominant features are various clay minerals alteromorphs after biotites and feldspars. The micromass is dominated by neoformation of Fe-oxyhydroxides, jarosite and clay minerals. The oxidation of sulfides and subsequent Fe-migration are evidence of the oxidized state of the surficial part of the waste rock piles.

Vegetation and a constant water regime induce rapid alteration of feldspar and delamination of biotite (Wongfun et al., 2013) associated with preferential leaching on the edge of the minerals. We observe that lichen frequently grows on the waste rocks here investigated. Entangled with the minerals of the weathered outer rim, lichen weakens the surface of the boulders (Fig. 3b). Roots of birch trees and ferns invading surficial fractures induce mechanical alteration and favor the development of protosoils. Therefore incipient weathering is increased in the vegetated piles.

3.1.2. Geochemical trends

The geochemical trends observed during incipient and further stages of weathering will be discussed as a function of the degree of chemical weathering, using the Chemical Index of Alteration (Nesbitt and Young, 1989), calculated as:

$$CIA = \frac{\% Al_2 O_3}{\% Al_2 O_3 + \% CaO + \% NaO + \% K_2 O}^* 100$$

CIA is widely applied to evaluate the chemical weathering intensity in weathering profiles and is used here to quantify the relative weathering intensity of our lamprophyre and granite samples, starting from typical CIA values between 55 and 65 for the least altered granites (Table S3 in the Supplementary material) up to the "optimum weathered value" of 100 (Price and Velbel, 2003). However, the Ca enrichment related to low temperature hydrothermal alteration precludes the comparison between weathered and hydrothermally altered granites. The most probable source of Ca is the convection circulation of meteoric waters induced by low temperature hydrothermalism. We have plotted the geochemical evolution of the waste rock piles in the $[Al_2O_3 - K_2O - (CaO + Na_2O)]$ diagram (Fig. 4) (Rollinson, 2014). Because of their high Fe concentration, the samples from "ferrihydrite pools" were excluded from this analysis. Two main trends of weathering appear; one from an incipient to an intense weathering state (trend A), and a second from an intense weathering state towards a pure kaolinite pole (trend B). The geochemical evolution of the four groups of rocks will be discussed by combining CIA values with the trends observed in the $[Al_2O_3 - K_2O - (CaO + Na_2O)]$ diagram.

Granites: most samples composing the waste rock piles are two-mica granites, with various alteration states. The mean CIA value is around 60 for unweathered/incipiently weathered granites, which means that no significant element loss occurs in the granitic rocks suffering incipient weathering. Values from 70 to 75 indicate weathered granites (Table S3 in the Supplementary material). In the ternary diagram, granite samples are grouped around 60–65 $Al_2O_3\%$, 30–35 K_2O % and 20–25% CaO + $Na_2O\%$ (Fig. 4). Two trends of weathering occur, the first one correlated with CaO + Na_2O leaching (trend A) and the second with K_2O leaching (trend B) essentially attributed to biotite and feldspar alteration. The granites mostly affected by trend B have the highest CIA values, 73 to 76 (Table S3 in the Supplementary material), referring to a probable supergene contribution to the alteration state.

Lamprophyres: the CIA value of the lower part of the lamprophyre dyke is 61, similar to a typical granite value. Highly weathered surficial lamprophyre dyke samples follow the "intense weathering" trend B



Fig. 4. Ternary Al_2O_3 - K_2O -(CaO + Na_2O) diagram used as a proxy to the alteration state of samples. Trend A goes from an incipient to an advanced weathering state. Trend B goes from an advanced weathering state to an intense weathering state towards a theoretical kaolinite pole (Al_2O_3).

towards an Al_2O_3 /kaolinite enrichment. CIA values for the surficial lamprophyre samples range from 68 to 77.

Hydrothermally altered samples: most CIA values for sulfide-rich samples are similar to the mean granitic values, ranging from 57 to 65. The three calcite-rich hydrothermally altered granites show low CIA values (from 36 to 49) (Table S3 in the Supplementary material). However, these CIA values are biased due to their high content of CaO (13 to 16%). CIA values of the calcite-rich and of the sulfide-rich samples are shifted towards the CaO apex or the K₂O apex respectively on the ternary diagram. Both groups follow the same trends of weathering (A and B) with an offset related to their mineral paragenesis. Within the calcite-rich group, the arrow on Fig. 4 points towards the sample with kaolinite with an important loss of CaO and Na₂O (Table S3 in the Supplementary material) suggesting an intense leaching.

Weathering products: CIA values of protosoils and technosoils (62–64) highlight incipient weathering in the coarse groundmass and immature technosoils. The humic fraction is extremely limited, correlated with L.O.I. values between 2 and 4%. More mature technosoils have CIA values between 71 and 73 correlated with higher L.O.I. values between 12 and 16% (Table S3 in the Supplementary material). This could be explained by mature overburden with higher OM content.

Regardless of the type of samples, the composition of the clay fraction is shifted towards Al_2O_3 when compared to the bulk samples, following trend A and/or trend B on Fig. 4. CIA leads to an overestimation of the alteration state if the samples were affected by hydrothermal alteration (Scarciglia et al., 2016). On the contrary, CIA values are underestimated in the calcite-rich (and to a lesser extent the sulfide-rich) samples. But as a whole, the use of CIA on the great variety of samples combined with the $[Al_2O_3 - K_2O - (CaO + Na_2O)]$ diagram demonstrates coherent weathering trends from granitic blocks to technosoil samples despite their chaotic setting in the waste rock piles.

3.2. Uranium speciation

3.2.1. Uranium minerals

Uranium bearing minerals were investigated with optical microscopy and SEM-EDS in the different groups of samples. Neither ore minerals nor other U bearing minerals may be identified by X-ray diffraction because of their minor occurrence in the rocks. However, microscopic investigations of the four groups of samples show the presence of accessory primary minerals, monazite and zircon (bearing up to 1% of U), with secondary U(VI)-minerals:

- 1 From the surface to the core, granitic rocks are affected by incipient weathering. Some isolated micrometric rosettes of U-oxyhydroxides (BEL-08-07, 315 ppm U) grow on smectite laths (Fig. 5a). These tabular-shaped U-oxyhydroxides are identified as schoepite $((UO_2)_8O_2(OH)_{12}$ ·12(H₂O)). 20 to 40 µm lamellar autunites (Ca $(UO_2)_2(PO_4)_2$ ·10–12H₂O) precipitates on flat apatite surface (MAC-08-01, 100 ppm U) (Fig. 5b). In addition, apatite shows dissolution figures.
- 2 In the deep lamprophyre dyke (BEL-05-15, 15 ppm U), occasional fluorite and apatite, known to occur as primary phases in these rocks (Rock, 1984), were observed but no secondary U bearing mineral was spotted. By contrast, the highly weathered surficial lamprophyre dyke (BEL-05-08 to BEL-05-10, 221 to 493 ppm U) concentrates U. The only secondary U bearing mineral present is autunite as large layered crystals (Fig. 5f) presenting characteristic tabular sheets that grow in a clayey matrix.
- 3 In the sulfide-rich yellowish granite (FAN-01-01, 348 ppm U), submicrometric needles of uranyl sulfate minerals, associated to Th rich nanoparticles (Appendix C-d), crystallize within a jarosite and smectite assemblage (Fig. 5d). These rocks are partially covered by a 1 mm hard thick greyish crust. This crust is composed of a tubular

La-Ce uranyl phosphate mat with smectite sheets and lichen (Fig. 5c). In the altered granite FAN-02-C1C, with a higher U concentration (997 ppm U), secondary uranyl phosphates associated to Fe-oxides precipitate as small, ill shaped phases in quartz fissures (Fig. 6). Nanometric uraninites are trapped in large (> $100 \,\mu m$) pyrite crystals with etched surface shown by dissolution features (Appendix C-e). Bordering the pyrite crystal (Appendix C-f), secondary micrometric, ill shaped Fe sulfates and uranyl phosphates fill a fissure, suggesting oxidizing conditions destabilizing the pyrite, confirmed by local Fe and S concentrations. In the calcite-rich hydrothermally altered granites (PEN-08-11, 1500 ppm U), stellate needle aggregates of uranophane ($Ca(UO_2)_2Si_2O_7 \cdot 6H_2O$) cover the pitted surface of large calcite crystals, pointing out a dissolution process (Fig. 5e). U rich coatings, several ten micrometer large, are observed on quartz confirming U mobility in these samples (Appendix C-c).

4 Among the weathering products, secondary U minerals are rare in the arena. Patches of sub-micrometric platy curite $(Pb_3(UO_2)_8O_8(OH)_6\cdot3(H_2O))$ are encrusted in a smectite matrix (BEL-08-09, 3000 ppm U) (Appendix C-a, and b). Protosoils (PEN-05-08, 38 ppm U and PEN-05-11, 69 ppm U) reveal late jarosite rosettes growing on the surface of feldspars, quartz and various clay minerals (kaolinite, smectite and chlorite). Apart from primary accessory minerals, monazites and zircons, no U bearing minerals were observed in protosoils.

3.2.2. Uranium geochemistry

Of the 54 samples analyzed, 74% have an U content below the initial U "cut-off value" (300 ppm), among which 22% have U concentrations below the regional background value of 20 ppm (Table S3 in the Supplementary material). Most granitic samples are affected by incipient weathering (CIA values of 60 to 65), except for the subgroup of intensely weathered granite (CIA 75) and the surficial lamprophyre samples, as discussed above. Because of their specificity humic horizon sample and ferrihydrite-rich samples are excluded from plots but will be discussed later. The bivariate plot CIA/U indicates that weathering induces a reconcentration of U into the clay fraction (Fig. 7). Two domains are highlighted: the U geological background and the U "cut-off range". The first with U < 20 ppm, contains mainly granitic bulk rocks affected by incipient weathering and the subgroup with CIA values around 75, previously described as intensely weathered samples (§ 3.1.2). Only two granitic samples with low CIA values (around 60) have U concentrations above the initial cut off value (> 300 ppm), with 314 and 1000 ppm for samples BEL-08-07 and PEN-01-06, respectively (Table S3 in the Supplementary material). This exceptional U concentration of 1000 ppm may be explained by the sorting between ore and waste rocks, based on large volumes (§ 2.1). The second domain concerns the weathering products, the weathered lamprophyres and the hydrothermally altered, sulfide-rich granites. U enrichment is noticeable in the weathering products, increasing progressively from 30 up to 4000 ppm in the technosoils.

The highest U concentrations in the clay fractions suggest local U mobility and reconcentration in the finest granulometric fraction during arenization. The reconcentration of U in the clay fraction varies from 28 to 85% (Table S4 in Supplementary material). A similar trend for rocks affected by previous hydrothermalism is noticeable with an offset at the origin related to their higher bulk rock U concentration.

Phosphate minerals and Fe-oxides are common U(VI) sinks. Samples with about 1% Fe_2O_3 present U concentrations ranging from 2 to 50 ppm. When Fe_2O_3 content reaches about 2%, a higher (× 10) U concentration (100 to 1000 ppm) is observed in granites and hydrothermally altered samples (Fig. 8). This U enrichment is especially visible in most technosoils with U values within, or above, the "initial cut-off range".

The most common U bearing minerals are monazite and various



Fig. 5. SEM pictures of rock chips: (a) Secondary electron image. Nanometric rosettes of uranium oxyhydroxyde, schoepite (Sch), concentrated on a smectite (Sme) background, in granite (BEL-08-07). (b) Backscattered electron image of a granitic sample (MAC-08-01). Lamellar autunites (Aut) precipitate on apatite (Apt). Mica (M) and zircon (Zr) are present in the background. (c) Secondary electron image, hydrothermally in altered granite with sulfides (FAN-01-01) showing micrometric tubular La-Ce uranyl phosphate (Lan P) mat with smectite (Sme) sheets. (d) Backscattered electron image in hydrothermally altered granite with sulfides (FAN-01-01), showing sub-micrometric needles of uranvl sulfate minerals, associated to high local Th concentrations (U-sulfate + Th), in a jarosite (Jrs) and smectite assemblage. Mica is also identified in the background. (e) Backscattered electron image in hydrothermally altered sample (PEN-08-11) showing stellate micrometric needle aggregates of uranophane (Urp) covering the etched surface of calcite (Cal). (f) Backscattered electron image. Upper part of lamprophyre dyke (BEL-05-08) showing well crystallized layered autunites (Aut) in a clay minerals background.

secondary uranyl phosphates described above. Fig. 9 represents a P₂O₅-U bivariate plot. In incipient weathered granites and weathering products, P₂O₅ is between 0.3 and 0.4%. These values can be attributed to primary and/or secondary minerals. The low P₂O₅ concentrations (< 0.3%) in weathered granite, lamprophyres and calcite-rich hydrothermally altered granites are attributed to primary mineral leaching. The clay fractions in arena, technosoils and lamprophyres show a high enrichment in P₂O₅, up to 0.9%, with the exception of the calcite-rich samples in which uranophane was observed, where it stays constant. Moreover the slope of increase of U and P₂O₅ in the clay fraction compared to the bulk samples is always positive but varies widely between 0.7 and 8.2. Nevertheless, the highest P₂O₅ and U values in the clay fraction suggest neoformation of secondary uranyl phosphates and/or mechanical weathering with concentration of heavy minerals.

The high U concentrations might be explained by a mechanical concentration of primary accessory minerals. Petrological observations suggest that Zr and Ce concentrations can be used to assess the zircon and monazite content, respectively. Microprobe measurements have shown that these accessory minerals contain up to 1% of UO₂. In the blocks and arena, Zr and Ce concentrations increase in clay fractions relative to bulk samples, with the exception of the protosoil sample in which Ce decreases in the clay fraction (Table S4 in Supplementary material). To assess the correlation between mechanical weathering and geochemical enrichment, granulometric separations were performed on 7 samples from the weathering products group (2 arenas and 5 technosoils) (Table S5 in Supplementary material). Mineralogical and geochemical data for each fraction are compared for a representative technosoil (FAN-02-C2, 237 ppm U) (Fig. 10). The concentration of clay

minerals (chlorite and kaolinite) increases from gravel to silt/clay fraction, while that of primary minerals (micas and feldspars) decreases (Fig. 10a). The U concentration in the silt/clay fraction increases by up to one order of magnitude, compared to the gravel fraction. Within this granulometric distribution, U is well correlated to Fe₂O₃, P₂O₅, Zr and Ce and to the CIA (Fig. 10b to f). The best correlations observed are between U and CIA ($r^2 = 0.98$; n = 5), U and Fe₂O₃ ($r^2 = 0.89$; n = 5) and U and P₂O₅ ($r^2 = 0.94$; n = 5). Concerning primary accessory minerals, U and Zr concentrations are well correlated ($r^2 = 0.80$; n = 5) (Fig. 10e) whereas U vs. Ce gives a moderate correlation ($r^2 = 0.63$; n = 5) (Fig. 10c) implying U is preferentially concentrated in zircons rather than monazites.

4. Discussion

4.1. Site characteristics and differences

Sites stand out in geochemical trends only when a group was found in one particular location such as the hydrothermally altered granites with sulfides, only sampled on Fanay, the hydrothermally altered group with calcite only found on Pény, or the ferrihydrite pools only sampled on Hyverneresse. Therefore, separation of samples by group overrules the sampling site. In a time span as short as sixty years, no difference is observed between thirty-year or sixty-year-old sites.

4.2. Identification of incipient weathering

To determine the importance of weathering in a time span of sixty



Fig. 6. SEM backscattered electrons image and EDS mapping (U, P and Fe) in sample from sulfide-rich group (FAN-02-C1C). Fissure in quartz filled with uranyl phosphate and Fe oxide.

years, this study focused on the surficial part the piles. Field observations have been combined with geochemical, mineralogical and micromorphological analyses to identify the various alteration processes in the waste rock piles. Meteoric water circulation and vegetation penetration induced chemical alteration and mechanical constraints in most surficial granitic blocks (20 cm large on average). Arenization of the blocks is more efficient if the blocks are affected by previous alteration (hydrothermalism, supergene). On the arena, protosoils appear with the development of lichen or the growth of birch trees.



Fig. 7. Bivariate plot showing the evolution of CIA values with log U concentrations. The same colors and symbols are used as in Fig. 8. Blue and red frames were added to indicate, respectively, the Uranium concentration corresponding to the geological background (2 to 20 ppm) and the uranium concentration corresponding to the initial cut-off value (100 to 300 ppm). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)



Fig. 8. Bivariate plot showing evolution of the $\% Fe_2O_3$ with U (ppm). The same colors and symbols are used as in Fig. 8. Bulk technosoil samples are indicated by arrows.

Weathering intensity, quantified by CIA, is assumed to be incipient when 55 < CIA < 65 and intense when CIA > 70. Therefore, the 4 granitic samples, out of the 27 investigated, which have CIA > 70, were probably affected by a supergene alteration episode prior to the waste rock pile edification. Indeed, similar CIA values have been observed on natural regoliths in temperate climates (Nesbitt and Young, 1989; Price and Velbel, 2003; Scarciglia et al., 2016). As pointed out above, samples affected by previous hydrothermal episodes should be handled with care in such geochemical diagrams since hydrothermalism can modify CIA values (Scarciglia et al., 2016). Therefore, such geochemical classification must be complemented by a thorough mineralogical identification.

The trends (A and B, Fig. 4) are characteristic of a loss of CaO, Na₂O and/or K₂O during weathering and thus, of an Al₂O₃ enrichment. This is concordant with an increasing concentration of secondary clay mineral. As a general conclusion, geochemistry data are consistent with the fact that the main weathering products are clay minerals (kaolinite, chlorite, smectite and interstratified clay minerals), phosphates and Feoxides.

In order to assess the effect of long term weathering of these waste rock piles, lamprophyre outcrops were analyzed for mineralogical and geochemical signatures. The good fit of deep lamprophyre rock to



Fig. 9. Evolution of $\% P_2 O_5$ with U (ppm). The same colors and symbols are used as in Fig. 8.

incipient weathering pole and surficial lamprophyres to highly weathered samples, in the overall geochemical trends, indicates that using surficial lamprophyre as a weathering proxy could provide a model for assessing the waste rock behavior under long periods of weathering.

4.3. Uranium bearing minerals

No remnant of U ore minerals (uraninite, coffinite, pitchblende) has been encountered in the waste rock-pile. Similarly, previous weathering studies on the Saint-Sylvestre two-mica granitic massif have shown the absence of uraninite in the first 50 m (Barbier and Ranchin, 1969). The absence of uraninite in our samples, when samples contain > 30 ppm of U, is an indicator of the oxidized state of the surficial waste rock pile under leaching conditions. When U concentration is below 30 ppm samples might never have contained uraninite, this concentration being explained by the presence of zircon and monazite.

Because the secondary bearing minerals crystallize in the bulk of surficial rock it is unlikely that U was transported by percolating water in the rock fractures. Therefore the absence of primary U(IV) is most likely due to oxidation of primary U(IV) minerals. Although we did not observe micron size uraninites trapped in quartz or feldspar, like in the deeper part of the piles (Boekhout et al., 2015), nanometric uraninites were found trapped in large pyrites suggesting relict reduced areas in the surficial pile (Appendix C-e). As pyrite dissolution would ensue in an oxidizing environment, uraninites would be dissolved, U(IV) oxidized to U(VI) and secondary U minerals precipitated.

Monazites and zircons are the only U bearing primary minerals present in all samples with a maximum U concentration of 1%. However, these minerals are resistant to weathering (Scherrer et al., 2000) and cannot be considered as a source of U during incipient chemical weathering. Mechanical weathering concentrates zircons and monazites in the clay fraction, due to their small size (1–20 μ m). Considering the U concentration in zircon and monazite and the Zr and Ce mean concentration in granites, these phases cannot account for > 10 to 30 ppm U in the bulk samples.

Secondary U bearing minerals crystallize in different environments controlled by recent processes and affected by geological history: in calcite-rich sample uranophane precipitates on a calcite surface. Calcite is known to be an efficient uranyl sorbent (Dong et al., 2005). Therefore precipitation of uranophane can be interpreted as a two steps mechanism of U trapping by sorption and further precipitation of uranophane concordant to calcite dissolution. Uranyl silicate precipitation as coating on quartz (Appendix C-c) is also experimentally observed with SiO₂ colloids and was shown to happen at slightly acidic pH (Lamb et al., 2016) comparable to the pH range of waste rock piles.



Fig. 10. Combined diffractograms and chemistry analyses results for the different granulometric fractions of one technosoil sample (FAN-02-C2). (a) DRX of different grain sizes, for the same technosoil sample, showing evolution of clay content from the gravel to the silt/clay fraction. Abbreviations translate as follows, Chl: chlorite, Ilt-Vrm: illitevermiculite, Bt: biotite, Ms.: muscovite, Kln; kaolinite, Qz: quartz, Kfs: K-feldspar, Pl: plagioclase, Px: pyroxene, FeOx: Fe oxide. (b) through (f) granulometric separation chemical analysis plots of U against Fe₂O₃, P₂O₅, Zr and Ce and CIA values.

In sulfide-rich samples, La-Ca-Th-U-P tubules in the external greyish alteration crust are comparable to florencite/rhabdophane precipitation after apatite weathering (Taunton et al., 2000). In the bulk of the sample uranyl sulfates co-precipitates with hydrated hexavalent U or Th oxides is possibly related to weathering of the uraninite-thorianite solid solution series. In the arena sample, curite is mixed with clay minerals (Appendix C-a and b). Curite is known to enhance the formation of insoluble uranyl phosphates (Finch and Ewing, 1992). Crystallization of various secondary U minerals in the different groups of samples proves the U trapping depends on the geological history of the sample. This has an impact on U mobility. Uranyl phosphates, such as meta-autunite, are known in natural systems as U sequestration products during weathering (Barbier and Ranchin, 1969; Jerden and Sinha, 2003; Jerden et al., 2003; Jerden and Sinha, 2006; Abd El-Naby and Dawood, 2008). In most granitic samples, lamellar autunites crystallize and form thin coatings on apatite (Fig. 5b). Their micromorphology and presence is typical of incipient weathering. The different morphology in the highly weathered lamprophyre (autunite layered crystals (Fig. 5f) with tabular sheets in the clayey matrix) suggests a more mature process of supergene alteration. Therefore, the autunites in most granite from the piles are formed from incipient weathering, whereas autunite growth in lamprophyre is formed over a long period of supergene alteration. The regular occurrence of uranyl phosphates in diverse samples from the surface of the piles confirms a process of U sequestration in specific minerals. Given the P2O5 concentration in samples uranyl phosphates could be the main U trapping process for U concentrations up to 4550 ppm (Table S3 in the Supplementary material). Still, not all P₂O₅ is available for U sequestration, a significant part belongs to apatite, monazite and other primary phosphates, and therefore other sequestration processes must also be considered.

4.4. Overall uranium speciation in the waste rock piles

Ferrihydrite pools and technosoil downslopes concentrate weathering products. No secondary U minerals were identified in ferrihydrite samples. Nevertheless, the high U concentration (2500 ppm) combined with the significant amount of P_2O_5 (2.5%, i.e. five to ten times more than in other samples from the waste rock piles), suggests an intricate process in U sequestration. Ferrihydrite is known to have an important role in U sequestration by sorption processes (Waite et al., 1994; Reich et al., 1998). The role of phosphate in enhancing U adsorption on ferrihydrite was evidenced and attributed to the formation of ternary complexes involving UO_2^{2+} and PO_4^{3-} (Payne et al., 1996). In the ferrihydrite samples of the pools the slightly acidic pH favors these ternary complexes. Trapping of important amounts of U by ferrihydrite suggests either chemical migration of U towards downslope weathering products, or mechanical migration of micrometric secondary uranyl phosphates and chemical reactivity in the ferrihydrite pool.

More mature technosoils bear the highest U concentration; with eventually an L.O.I. value of 48.5% which indicates a high organic matter content, highlighting the important influence of organic matter in U sequestration (Cumberland et al., 2016). U sequestration by OM should be considered as a long term sink with development of vegetation on waste rock piles. Nevertheless in most samples of the piles, OM is negligible and other U sinks may be responsible for U sequestration.

In clay fractions, U enrichment is observed in lamprophyres with interstratified clay minerals and in granitic arenization matrix and protosoils with smectite. U could be sequestrated by clay minerals (Tsunashima et al., 1981; Chisholm-Brause et al., 1994; Schlegel and Descostes, 2009). The sorption efficiency of smectite favors the nucleation of uranyl minerals (Schindler et al., 2015), limiting U migration in the pile. Clay minerals are known as an efficient U sequestration system, they are generally used as a liner to limit environmental impact of metallic pollutants (Rowe et al., 2004). Fe-oxides and clay minerals

play an important role in radionuclide migration, efficiently trapping uranyl as inner- and/or outer-sphere complexes at their surface (Brown and Calas, 2012; Neiva et al., 2014; Cabral Pinto et al., 2014).

Granulometric separations have shown a good correlation of clay minerals, P_2O_5 and Fe-oxides to U concentration. The sorption efficiency of the major phases (clay minerals, Fe-oxides) is probably the dominant process of U sequestration; however, the presence of nanometric U bearing minerals embedded in these divided secondary phases should not exclude the possible role of phosphates in efficient sequestration.

5. Conclusion

No trace of relict uranium ore was found in the investigated samples. The scarcity of uraninite trapped in pyrite confirms the global oxidation state of the surficial waste rock pile. The weathering degree of each sample, in the chaotic deposits of waste rock piles, was successfully assessed relying on the CIA value (around 60 for incipient weathering, and 70 for intense weathering), the mineralogy and the micromorphological observations. Hexavalent uranium is known for its high mobility, hence its early migration within the waste rock pile since the beginning of the incipient weathering. It is concentrated in the finest fraction, the U content being controlled by the low solubility of neoformed uranyl minerals, mainly uranyl phosphates, the cationic exchange capacity of smectites and the sorption efficiency of ferrihydrite. The evolution of oxidation in depth of the waste rock pile and the associated weathering processes were investigated in a preliminary study (Boekhout et al., 2015), showing overall oxidation of the waste rock pile with dominantly uranyl phosphate precipitation. Secondary uranyl phosphates are important in the presently investigated surficial samples, as they are the dominant secondary U mineral found and are known to cause low levels of aqueous uranium in groundwaters due to their low solubility (Munasinghe et al., 2015). The different micromorphology of autunites, from well-defined layered crystal in lamprophyres compared to isolated lamellae in granites, has been related to the duration of weathering. Other secondary uranyl minerals also ensure U trapping, such as uranophane and uranyl sulfates. The present investigation shows that large amounts of Fe oxyhydroxides concentrate downslope of the waste rock piles. This important result confirms the primary role of Fe-oxyhydroxides in limiting U mobility under slightly acidic pH domains and oxidizing environments. Based on the influence of surficial incipient weathering of the waste rock piles, a realistic assessment of the environmental evolution will imply investigating deep weathering processes, under different redox conditions.

Supplementary data to this article can be found online at http://dx. doi.org/10.1016/j.gexplo.2017.08.010.

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Appendix A. Regional geological background

The St-Sylvestre peraluminous leucogranitic complex was emplaced 324 Ma ago, as a part of the Variscan orogeny (Scaillet et al., 1996, and references therein). The most representative rocks are coarse to medium-grained, foliated, with U concentrations ranging from 10 to 30 ppm. Those two-mica granites underwent a complex hydrothermal alteration history, including the formation of episyenites (columnar structures related dominantly to quartz dissolution):

- From 295 to 285 Ma, lamprophyre intrusions crosscut the granitic bodies, during brittle fracture tectonic events (Rock, 1984). "Lamprophyre dykes [...] contain augite, Mg-rich biotite, feldspar (potassic and/or plagioclasic), pseudomorphs of minerals of the olivine group, apatite, and magnetite. In nearly all samples round quartz grain with a pyroxene rim can be observed. They are interpreted to be relicts of granitic material, partially assimilated by the lamprophyre magma (Velde, 1971)." (Leroy, 1978). Lamprophyre intrusions cause the fluids to heat in the granitic massif fractures inducing hydrothermal alteration.
- Hydrothermal activity starting at 345 °C induced the episyenitization of the granite, with formation of sulfides (pyrite, marcasite) and U ore mineralization (pitchblende) dated at 284–274 Ma. U enrichment occurs up to 10 m. from the ore vein (Ménager et al., 1992). The enriched zones are characterized by a gradient in sulfides and uranium ore minerals. Cooling of the system (from 330 °C to 140 °C), after pitchblende deposition, induced formation of coffinite, hematite and marcasite. At temperatures below 135 °C calcite, barite and fluorine minerals precipitated (Leroy, 1978).
- Recent supergene alteration starts around 30 Ma ago (Leroy, 1978). U-containing minerals are either related to the magmatic emplacement of the granite (mostly monazite, but also zircon and magmatic-stage uraninite), or to the hydrothermal mineralization (pitchblende and coffinite). Supergene alteration is thought to go as deep as 100 m. (Cathelineau, 1983) but is mainly located in the first 15 m. (Barbier and Ranchin, 1969). Products from supergene alteration are similar to intense weathering products, uraninite is dissolved and clay minerals, Fe oxides and autunites are formed.

Appendix B. Location of the former mining sites



Appendix C. Additional SEM data (a) Backscattered electron image in arenization matrix sample BEL-08-09, from Bellezane showing platy curite (Cur) crystals in clay mineral background. (b) Secondary electron image, close up from picture (a) platy shape of nanometric curite crystals is visible. (c) Secondary electron image, from hydrothermally altered granite PEN-08-11 showing U-coating on quartz (Qz) mineral. (d) Backscattered electron image of a similar mineral to the one in Fig. 5d, showing micrometric uranyl sulfates needles with Th local concentration (U-sulfate + Th). Submicrometric particles (in brighter color in the needles) are associated to the needles. (e) Backscattered electron image from sulfide rich sample (FAN-02-C1C), showing nanometric uraninites (Urn), circled in black, in pyrite (Py). (f) Close up from (e) showing micrometric U-phosphate (U-ph) and Fe-sulfate in a fissure of a quartz mineral. Elements added (Fe, S and U) are indicated because they were identified in the background



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