
Light in the cave: Opal coating detection by UV-light illumination and fluorescence in a rock art context

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Abstract:

The formation of silica coatings on the cave walls of the Points cave raises questions about the analytical access to the specificities of the pictorial material (geochemistry and petrography); and about the state of conservation of the rock art. Conventional *in situ* spectroscopic techniques (pXRF, pRaman) are ineffective to identify and characterize these coatings. In this study, we propose to use a UV fluorescence method for the detection and recognition of opaline coatings, based on the fluorescence specificities of the uranyl-silica complexes composing these deposits. Spectral identification using UV laser-induced fluorescence spectroscopy coupled with UV illumination was performed on samples, μ -samples and on the walls of the Grotte aux Points rock art site. The well-defined peaks observed in the fluorescence emission spectra due to uranyl ions validate the detection of the complex opal-uranyl and its correspondence with the green fluorescence observed under UV light at micro and macroscopic scales. *In situ* optical measurements under UV illumination reveal the presence of an opal layer, in particular on the rock art walls of the cave. Observations on the occurrence and distribution of opal provide the first insights into the evolution of the walls and the chronological constraints on the development of the opal layer. regarding the interactions between the silica coating and the pigment suggested by the multi-scale observations of the μ -samples, it opens the question of rock art conservation. Thus, by developing a specific method of non-destructive characterization of opal coatings, this study start a new approach for the study of the taphonomy of decorated walls and proposes to use siliceous mineralization both as a marker of the natural history of caves and as an index of their use by ancient human groups.

Keywords: silica coating; uranyl; UV fluorescence; *in situ* detection; cave art



1. Introduction

Natural activity in caves, mostly water weathering (Chalmin *et al.* 2019; Delvigne 1998; Salomon *et al.* 2021), transforms the surface of the walls by physical, chemical, biological and mechanical action. Thus, the traces of all these transformations (environmental input) can be a source of information regarding natural and anthropological events on the wall surface, such as drawing and painting realizations, cave environment evolution, and human presence in the cave (Monney & Jaillet 2019; Pons-Branchu *et al.* 2014; Quilès *et al.* 2015; Sadier 2013; Shao *et al.* 2017; Valladas *et al.* 2017;).

Among the taphonomic processes impacting rock art pictorial matter, mineral-coating formation as weathering products is well described in rock art research (Bassel 2017; Chalmin *et al.* 2018; Huntley 2012; Mauran *et al.* 2019; Vignaud *et al.* 2006). Silica rich amorphous deposits, also called silica skins, have been observed at different cave and open-air rock art sites (Aubert *et al.* 2004; Aubert *et al.* 2012; Huntley 2012; Watchman 1990). Those studies have suggested both a positive and negative impact on rock art conservation due to opal coating development. Indeed, the strong interaction suspected with hematite pigments has been suggested as an element of conservation enhancement, notably compared with other pigments (Watchman 1990). However, some authors have also observed exfoliation processes of silica skins, which could play a role in rock art weathering (Aubert *et al.* 2004; Green *et al.* 2017). To our knowledge, there is still no clear answer on the role of silica coatings as conservation factors of pictorial matter. In addition to this conservation issue, silica skins have been proposed as tools for indirect dating of rock art, especially in the case of inorganic pictorial matter (Aubert *et al.* 2004; 2012). Indeed, this mineral phase is known to be enriched in uranyl ions, but the U-Th dating application remains hypothetical due to silica skin thickness and absence of stratigraphy, which complicate both sampling and measurement reliability (Green *et al.* 2017).

Thus, silica skin characterization represents a key issue in the context of rock art studies. However, it remains difficult to identify and characterize, especially with non-invasive techniques. Currently, the use of *in situ* spectroscopic techniques in rock art studies is increasing, as these methods can provide information on both pictorial matter and pigment environments (substrate, deposits, concretions, *etc.*). The portability and decreasing cost of instruments coupled with the rapidity and the non-destructive character of analyses have led to a quasi-systematic use of these techniques in recent rock art studies (Chanteraud *et al.* 2021; Huntley 2012; Mauran *et al.* 2019; Trosseau *et al.* 2021). However, amorphous silica characterization, even in rock art context, is generally based on laboratory observations such as SEM or XRD analyses (Gaillou *et al.* 2008; Garcia-Guinea *et al.* 2013; Green *et al.* 2017; Huntley *et al.* 2015; Watchman 1990). In addition, the signal of pictorial matter acquired with portable spectroscopic techniques could be impacted by the presence of silica skins, as observed by Huntley (2012) in the case of pXRF measurements.

In this paper, we propose a new method for *in situ* detection and characterization of amorphous silica in a rock art context based on UV laser-induced fluorescence (LIF). Indeed, uranyl fluorescence characteristics under UV light are well known and have been observed in silica mineralization, especially in opal (Fritsch *et al.* 2001; Gaillou *et al.* 2008; Gorobets *et al.* 1977; de Neufville *et al.* 1981). UV spectroscopy presents the same advantages as other portable spectroscopic techniques, but the bright green fluorescence and the specific spectral features displayed by uranyl ions enable the targeted identification of opal coatings. To our knowledge, only one study has reported opal detection in caves using optical methods based on UV techniques (Garcia-Guinea *et al.* 2013), and no study has applied UV spectroscopic methods in a rock art context for opal identification. Here, we propose a methodological

development based on laboratory and field experiments to validate the use of *in situ* UV techniques for opal detection in a rock art context by coupling *in situ* optical and spectroscopic analyses to obtain multi-scale information.

This study was performed in Points Cave (Aiguèze, Gard, France), which contains an important spread of opal coating on the cave walls. In this approach, Points Cave perfectly illustrates the importance of environmental input characterization in the study of rock art (Chanteraud *et al.* 2021). In Points Cave, some pieces of the rock wall bearing colouring matter were discovered during archaeological excavations in front of some rock art panels which allowed transport, μ -sampling and analysis in the laboratory. Analysis of these coloured flakes allows us to identify and measure the environmental input, which modifies and obscures the identification of pictorial matter characteristics.

2. Materials

2.1. Study site: Points Cave

Located in the Ardèche River Canyon, less than 10 km downstream of Chauvet Cave, Points Cave is a Palaeolithic rock art site identified in 1993 (Figure 1) (Deschamps *et al.* 2018). Archaeological studies have been performed since 2011 as part of the “Datation Grottes Ornées” project (“Cave Art Dating” project) (Monney 2011; 2018). The entrance opens on a hundred-meter long cave in Urgonian limestone. The rock art which is exclusively composed of red drawings and paintings, is located in the middle part of the cave preserved from sunlight (Figure 1). Excavations conducted at the entrance indicated human and animal occupation since the late Pleistocene to the present day (Monney & Jaillet 2019).

Points Cave is currently disconnected from hydrogeological flows (Jaillet & Monney 2018). Only a few infiltrations can be observed after strong precipitation events. The low level of leaching on the wall and the low relative presence of a calcite veil are due to this weak hydrogeological activity. Millimetric to centimetric concretions and efflorescence’s (coralloid type of crystallization) have also developed on the wall surface in the rock art sector (Barbaran & Nouet 2020; Mauran 2019).

2.2. Points Cave rock art and pictorial matter

Rock art, composed exclusively of red drawings and paintings, is currently observed in the middle part of the cave. It comprises five animal figures (3 ibexes, 1 horse, 1 bison), five indeterminate tracings, two bi-lobed signs, one open-angle sign and four clusters containing a total of 59 palm-printed dots, commonly referred to as “palm dots” (Monney 2018a). On-site spectroradiometric analysis showed that Points Cave rock art is homogenous in colour (Lafon *et al.* 2022). Moreover, pieces of colouring matter were found during the archaeological excavations in the sedimentary sequence at the entrance of the cave (Chanteraud 2020: 62; Chanteraud *et al.* 2019).

Investigations carried out at the foot of the rock art panel known as the “Bi-lobed Signs” revealed the presence of five coloured wall flakes on the ground in the chaos of rocks: S-ECA-01 to S-ECA-05. These 3 to 20 cm flakes are composed of limestone fragments detached from the wall by mechanical action and covered on one side with colouring matter (Monney 2011). They more than likely come from the large bilobed sign (PTS-10) or from other unknown rock art entities that may have crumbled entirely from the walls nearby (Monney 2018b). These flakes are a major asset of Points Cave because they allow to investigate the cave walls surface with accurate instruments by bringing them in the laboratory.

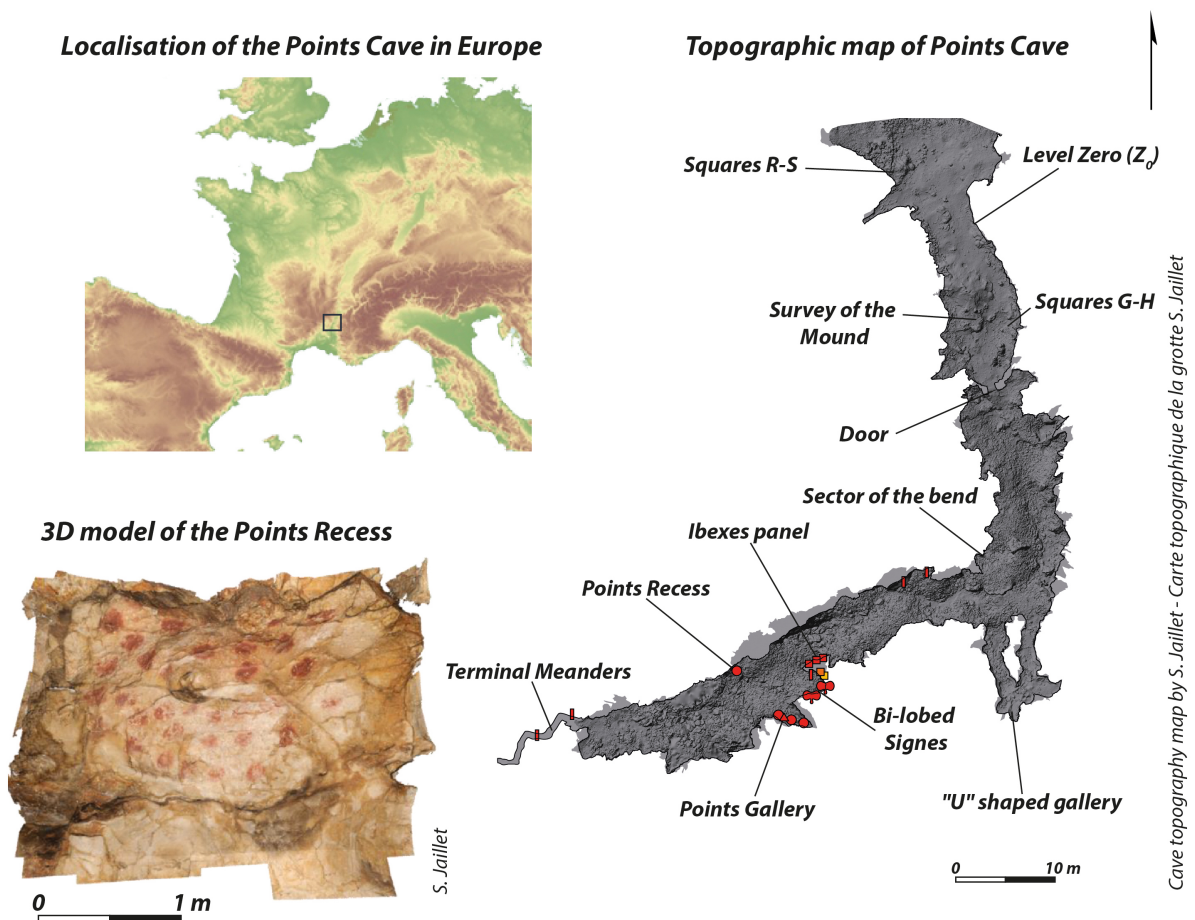


Figure 1. Points Cave location in southeastern France, topographic map of the cave with location of the excavations and the rock art panels (J. Monney); 3D model of the “Points Recess” panel (J. Monney).

Figure 1. Carte de localisation de la Grotte aux Points dans le sud est de la France, locus des sondages archéologique et des panneaux ornés dans la grotte (J. Monney), modèle 3D de de la « Niche aux Points » réalisé par photogrammétrie (S. Jailliet).

To identify both the iron-oxide morphologies and the natural mineralization that can form on the surface of the decorated walls, our study was focused on three of the five wall flakes (S-ECA-03, S-ECA-04, S-ECA-05) discovered in the rock art sector of Points Cave, because of the presence of opal coating on their surface (Figure 2). μ -sampling of pictorial matter was performed on the coloured flakes: μ -S-ECA-03, μ -S-ECA-04, μ -S-ECA-05. These last two have been fragmented during sampling giving: μ -S-ECA-04_prlvt_01, μ -S-ECA-04_prlvt_02 and μ -S-ECA-04_prlvt_03, and μ -S-ECA-05_prlvt_01 and μ -S-ECA-05_prlvt_02 (Supplementary Information1).

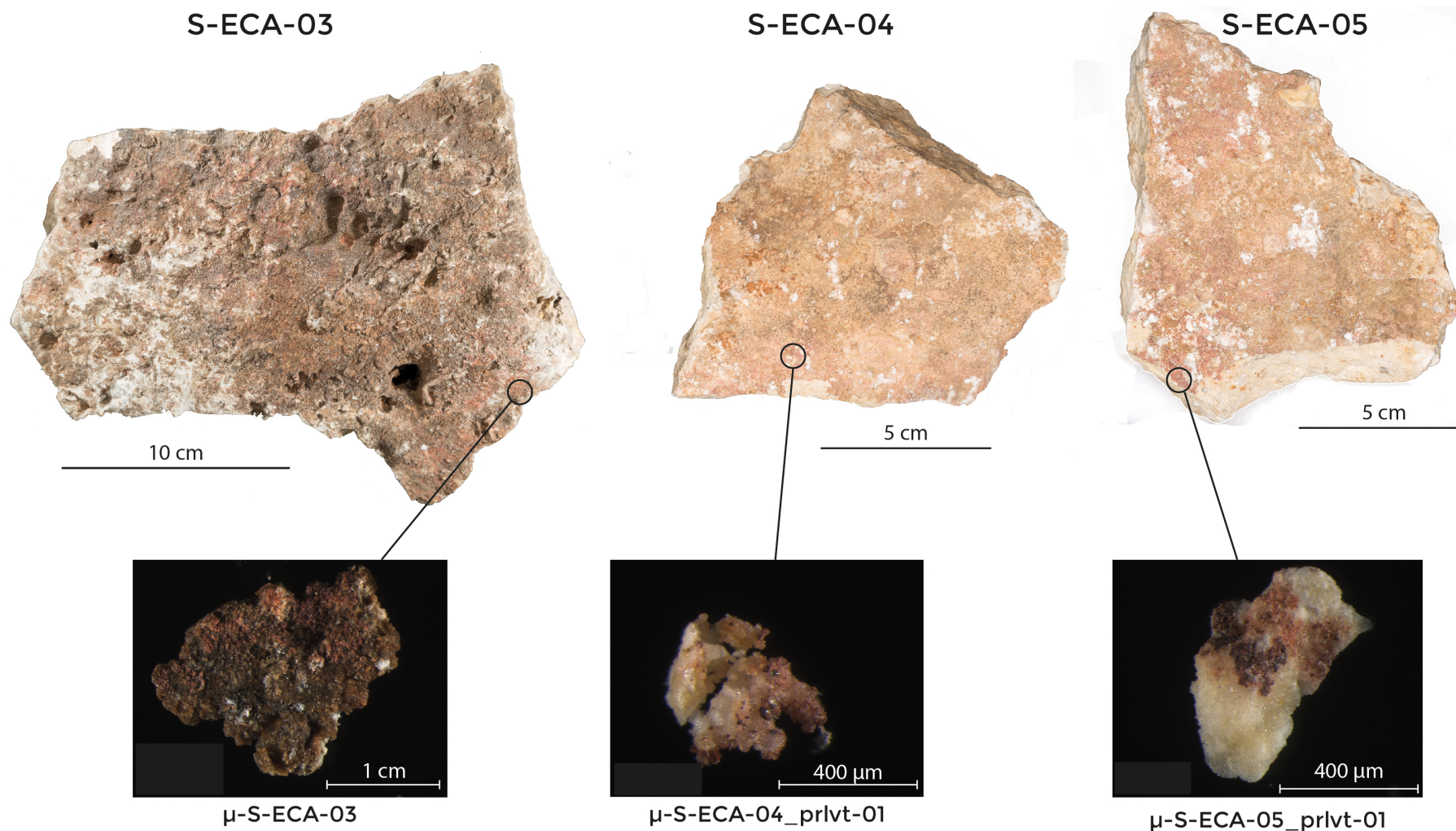


Figure 2. Limestone flakes found in the archaeological excavations at the foot of the Bi-lobed Signs panel and location of μ -sampling.

Figure 2. Ecaille de calcaire colorées détachées de la paroi retrouvées au pieds du panneau orné dit des signes Bilobés, avec localisation des μ -échantillons à la surface des écailles.

Moreover, 12 μ -samples of pictorial matter on limestone and 8 μ -samples of the undecorated cave walls (limestone) were directly sampled on and around the rock art panels of the cave and added to the study (Table 1, Supplementary Information 2). All μ -samples of pictorial matter were observed and analysed without any preparation.

Table 1. Inventory of μ -samples with colouring matter coming from the rock art panels.
Table 1. Inventaire des μ -échantillons réalisés sur les parois ornées de la Grotte aux Points

Sample N°	Rock Art	Type
- Ibexes Panel -		
Prm-19-01	Near to the Ibex n°5	Wall Blank
Prm-19-02	Ibex n°5	Pictorial matter
- Horse Panel -		
Prm-19-03	Near to the Horse	Wall Blank
Prm-19-04	Horse n°7	Pictorial matter
Prm-19-05	Bison n°8	Pictorial matter
- Bi-lobed Signs Panel -		
Prm-19-06	Near to the Sign n°9	Wall Blank
Prm-19-07	Small Bi-lobed Sign n°9	Pictorial matter
Prm-19-08	Large Bi-lobed Sign n°10	Pictorial matter
Prm-19-09	Palm-dot n°14-02	Pictorial matter
Prm-19-10	Open-angle Sign n°13	Pictorial matter
- Dots Gallery -		
Prm-19-11	Near to the Palm-dot n°11-02	Wall Blank
Prm-19-12	Palm-dot n°11-02	Pictorial matter
Prm-19-13	Palm-dot n°11-03	Pictorial matter
- Dots Gallery (cluster 11) -		
Prm-19-14	Near to the Palm-dot n°11-01	Wall Blank
Prm-19-15	Palm-dot n°11-01	Pictorial matter
Prm-19-16	Palm-dot n°15-21	Pictorial matter
Prm-19-17	Palm-dot n°15-01	Pictorial matter
- Points Recess -		
Prm-19-18	In the centre of the panel	Wall Blank
- Between Ibexes panel and Horse panel -		
Prm-19-19	On the limestone	Wall Blank
- Points Recess -		
Prm-19-20	Near to the Palm-dot	Wall Blank

3. Methods

3.1. Sub-section (part of the methods section)

3.1.1 Laboratory and in situ observations at macroscopic scale

Image capture for macroscopic observations was performed in two steps: 1) a photograph was taken under white light, and 2) another photograph was taken under UV light illumination. Observations at macroscopic scale in the laboratory and in the cave were realized with a Canon EOS 5D Mark III camera and a Canon EOS 7D camera fixed on a tripod. A detached flash was used for image capture under white light. For UV light

illumination, 4 UV LEDs (280 nm, 2.26 W, NewEnergy) were fixed on orientable macroflash bars on each side (2 LEDs per side) of the camera, allowing both macrophotography and general views of cave walls. Camera parameters are available in Table 2.

Table 2. Camera parameters used for both white and UV light illumination during field and laboratory macroscopic observations.

Table 2. Paramètres pour l'acquisition photographique des images sous illumination de lumière blanche et UV en laboratoire et dans la grotte.

			Aperture	Obturation speed (sec)	Iso
Laboratory	White light	data	F/7.1	1/320	1600
	UV light	data	F/7.1	10	1600
Field	White light	Macro	F/11	½	320
		Wall view	F/11	1/60	100
	UV light	Macro	F/4.5	½	320
		Wall view	F/11	15	320

Laboratory observations were also performed using a stereomicroscope (LEICA M165 C) under white and UV light. The same 4 LEDs were used for UV illumination.

3.1.2. Laboratory observations at microscopic scale

At microscopic scale, observations of μ -samples of flakes on carbon tape were realized with a field-emission scanning electronic microscope (SEM) ZEISS Ultra+ associated with an EDX (energy dispersive X-ray) probe (SDD, Bruker) working in high-vacuum mode with a 15 keV voltage. Images were taken with secondary electrons using in-lens or Everhart-Thornley detectors (SE2) and with backscatter electrons (BSE-AsB detector) (Néel Institut, Grenoble).

3.2. UV fluorescence analyses

Stationary fluorescence signal of flake and flake μ -samples was measured in the laboratory with a solid-phase spectrofluorimeter designed in the EDYTEM laboratory for non-destructive solid-phase measurements (Perrette *et al.* 2000).

This instrument is divided into excitation and detection compartments associated with a translation stage system for sample surface measurement. For this experiment, the excitation system was composed of a Nd:YAG laser (Crylas, FQSS266-Q1) with a 266 nm excitation wavelength. The fluorescence emission response was collected after sample excitation by focusing the laser beam ($\sim 30 \mu\text{m}$) on its surface. The detection system was composed of a low-pass filter monochromator (Jobin Yvon, MicroHr) for light diffraction, fitted with a 300-g/mm diffraction-grating centred at 620 nm, associated with a thermoelectric-cooled, back-illuminated CCD (Jobin Yvon, Sincerity) for high-efficiency signal detection in the UV-visible light domain. As no manipulation, modification or destruction of the sample surface occurred during analysis, this technique is suitable for archaeological sample measurements.

Two types of fluorescence measurements were realized:

Single point measurements were performed on flake μ -samples: μ -S-ECA-03, μ -S-ECA-04 and μ -S-ECA-05-02 (Supplementary Information 3). One spectrum was independently acquired on the sample. Measurement location was determined manually. An acquisition time of 1 s and a monochromator entrance slit of 0.25 mm ($\sim 1 \text{ nm}$ spectral resolution) were used for spot spectrum acquisition.

Surface measurement was performed on flake S-ECA-05 (Supplementary Information 4). The motorized translation stage system allowed movements in two directions for surface

measurements. The image was obtained by stacking lines along the Y-axis. For this study, fluorescence surface imaging was performed with a 100×100 µm resolution using a 0.1 s acquisition time and a 0.05 mm entrance slit.

Data pre-processing was performed with MATLAB software (Supplementary Information 5). Spectra were corrected from baseline and instrument responses and then filtered with the Savitsky-Golay method (Savitzky & Golay 1964).

4. Results

4.1. Macroscopic scale observations

At macroscopic scale (by visual inspection), beside the centimetric coralloid crystallization there is no other mineralization visible than the limestone on the walls, flakes or µ-sample surfaces. In fact, at this observation scale, surfaces seemed to be well preserved with strong colouration from the pictorial matter and clear access to the calcareous substrate (Figure 2).

Despite the results of the visual inspection of the walls and the observation of the µ-samples both of the rock art and the flakes under stereomicroscope, pXRF analyses (1 cm³ spot) showed that decorated panels of the cave display mineralization, such as calcium sulphate (Chanteraud *et al.* 2020; Supplementary Information 6). This contradiction between observation and geochemical analysis reveals the need for microscopic inspection using SEM device.

4.2. Microscopic scale observations

At the microscopic scale, a 1 µm-thin mineral film consisting of nonordered spheres was identified as silica mineralization on µ-S-ECA-03, µ-S-ECA-04-prlvt-02, µ-S-ECA-05-prlvt-01 (Figure 3 and Supplementary Information 2). However, µ-S-ECA-03 shows less opal coating than the two other µ-samples. The geochemistry and hummocky morphology of the opal could be related to a type A-g amorphous opal with a SiO₂*n*H₂O formula (Flörke *et al.*, 1991). The mineral structure present on coloured flakes suggests its formation from a water film containing a high concentration of silica (Curtis *et al.* 2019; Monger & Kelly 2002). With this mode of formation, it is difficult to assume if the opal came before the hematite deposit. In fact, it is possible that the opal coating grows at the interface between the limestone substrate and the colouring matter.

Concerning the µ-samples directly collected on the decorated walls, the same observations were noted, including pseudo-hexagonal platy hematite and strong indications of opal (Supplementary Information 1). Importantly, these samples showed significant opal development, with a complete coating of the pictorial matter to the extent that the morphology of the hematite was no longer observable on the surface (Figure 3). Weathered hematite plates seem to have been "ingested" by the silicate coating and could only be observed when a section was accessible on the surface of the sample (Figure 4 and Supplementary Information 2). However, preliminary studies on the walls of Points Cave using portable spectroscopic techniques (pXRF and Raman) could not identify the presence of opal coating *in situ* (Chanteraud *et al.* 2020).

μ -S-ECA-05-prlvt-01

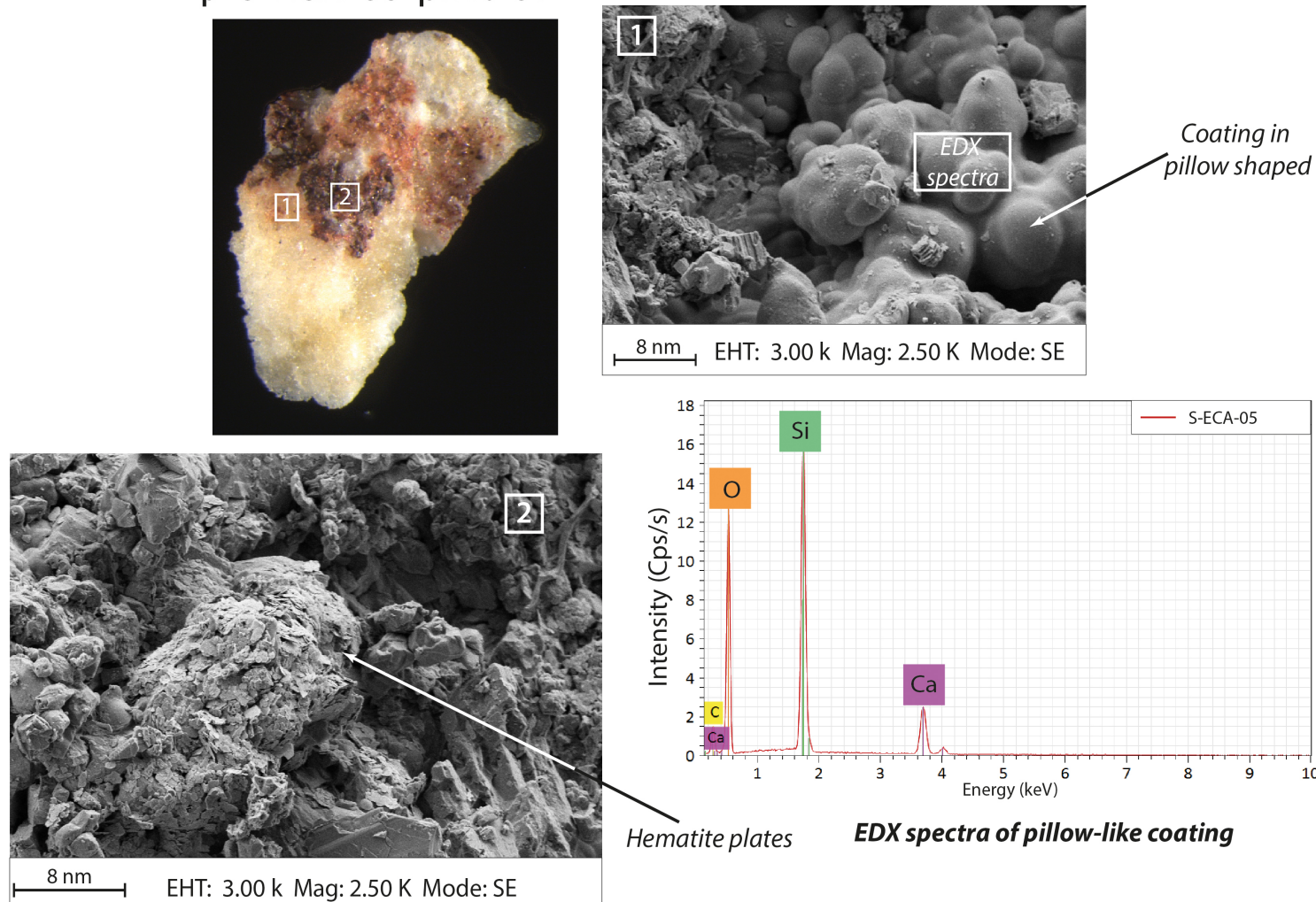


Figure 3. SEM observations of μ -S-ECA-05_prlvt_02. 1. Hematite plates observation in Secondary Electron mode (SE); 2. Pillow-like silica coating on surface in SE mode; White rectangle = Area of EDX spectra on the pillow-like silica coating.

Figure 3. Observations MEB des μ -prélèvement μ -S-ECA-05_prlvt_02. 1. Observation des plaquettes d'hématite en mode électrons secondaires (SE) ; 2. revêtement de silice en forme de coussin sur la surface en mode SE ; rectangle blanc = zone des spectres EDX sur le revêtement de silice en forme de coussin.

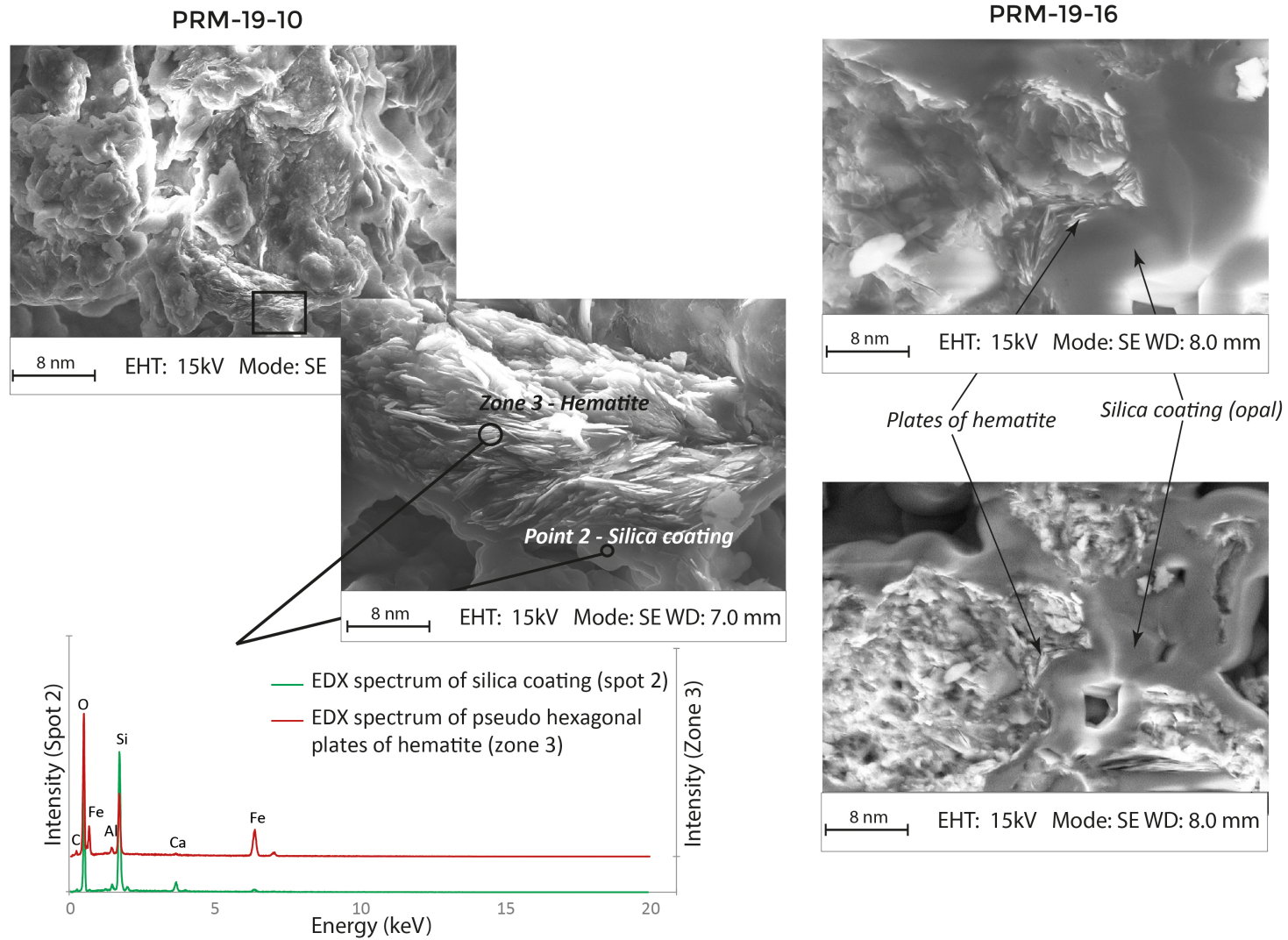


Figure 4. Hematite plates embedded in the silica coating on PRM-19-10 and PRM-19-16 μ -samples (SEM observation in SE and BSE mode).

Figure 4. Plaquettes d'hématite recouvertes par le revêtement siliceux sur les micro-échantillons PRM-19-10 et PRM-19-16 (observation au MEB en mode SE et BSE).

4.3. Opal identification by UV-fluorescence

Fluorescence analyses were first performed on 3 flake μ -samples (μ S-ECA-03, μ S-ECA-04 and μ S-ECA-05) that were previously characterized by different geochemical analyses and on which opal was detected. Due to the small size of the samples (approximately 200 μ m), only a few localized spectra were recorded. The spectra obtained could be divided into different emission regions depending on the main signal sources (Figure 5, Supplementary Information 3):

From 300 to 470 nm, this region corresponds to the fluorescence emission of the sample matrix. Peaks and shoulders were detected at approximately 350 nm, 380-390 nm, 410 nm and 430-440 nm. According to the literature, they could be associated with organic matter entrapped in the crystalline matrix (Van Beynen *et al.* 2001; McGarry & Baker 2000; Perrette *et al.* 2000; Perrette *et al.* 2005; Quiers *et al.* 2015) or with the silica material fluorescence response to UV-light excitation (Boyko *et al.* 2011; Garcia-Guinea *et al.* 2013).

From 470 to 750 nm, special features were identified in this part of the spectra for samples μ S-ECA-04 and μ S-ECA-05. They were characterized by a sequence of 3 defined peaks at 501, 521, and 545 nm and a shoulder at approximately 572 nm. These peaks were identical in all spectra measured on μ -samples μ S-ECA-04 and μ S-ECA-05 and coincided with the uranyl ion spectrum in silica matrices based on a review in the literature (Table 3), but were absent from μ S-ECA-03 spectra.

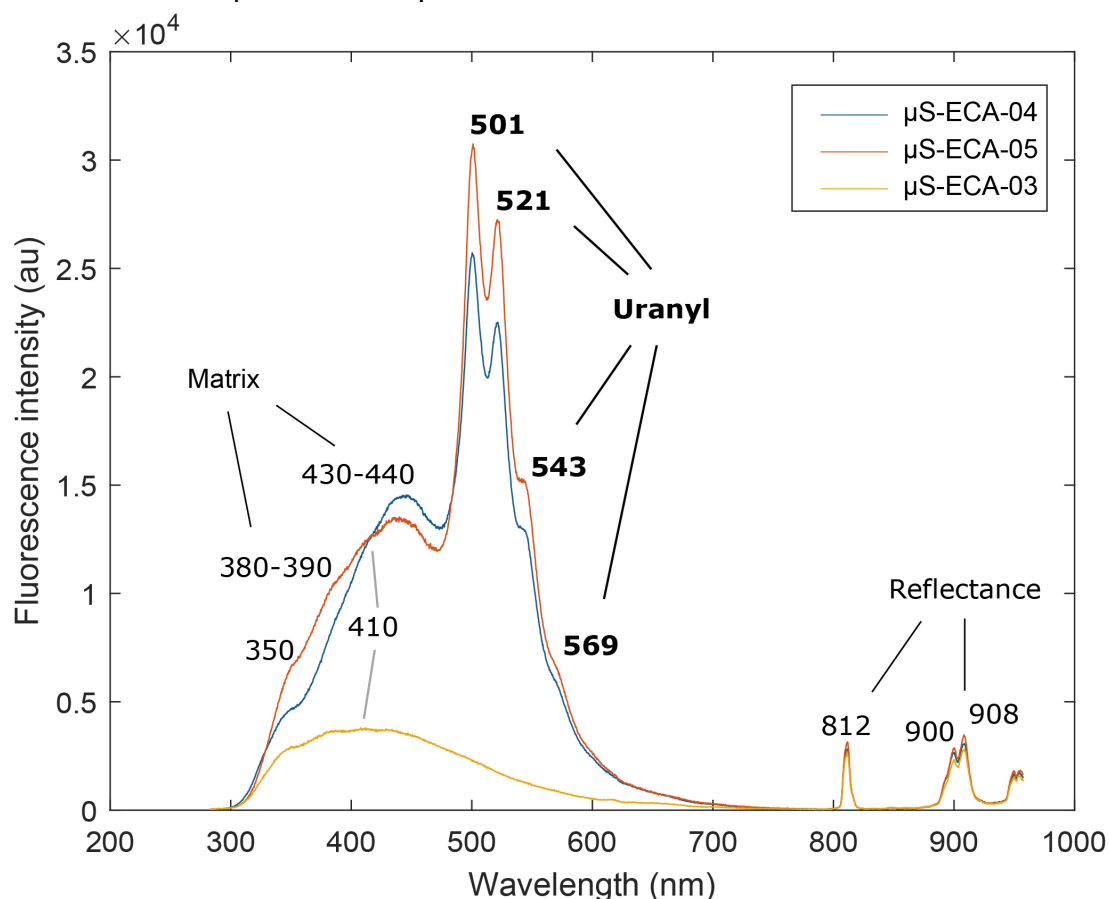


Figure 5. Mean fluorescence emission spectra (excitation: 266 nm) of samples μ S-ECA-04, μ S-ECA-05 and μ S-ECA-03. Spectra are divided into three different regions as a function of the main fluorescence signal sources: sample matrix, uranyl ions and laser emission reflectance.

Figure 5. Spectres d'émission de fluorescence moyens (excitation : 266 nm) des échantillons μ S-ECA-04, μ S-ECA-05 et μ S-ECA-03. Les spectres sont divisés en trois régions différentes en fonction des principales sources de signaux de fluorescence : matrice de l'échantillon, ions uranyle et réflectance de l'émission laser.

Entrapment of uranyl ions in siliceous matrices, especially opal phases, is well documented in the literature (Devès *et al.* 2012; Fritsch *et al.* 2001; 2015; Gaillou *et al.* 2008; Kasdan *et al.* 1981; Kinnunen & Ikonen 1991; Neymark *et al.* 2000; Othmane *et al.* 2016; Zielinski, 1980). The strong affinity of uranyl groups for amorphous silica leads to a strong U-opal association, which is stable at the scale of geological time (Othmane *et al.* 2016). As opal was the only silica phase identified on these samples, uranyl-specific spectra could be associated, in this case, with the presence of opal on samples, and UV fluorescence analysis represents an efficient tool for its identification.

As opal detection was subject to uranyl fluorescence properties, detection using UV fluorescence was dependent on uranium entrapment in silica crystalline structures. In their study of opal gems from different geographic and geological contexts, Gaillou *et al.* (2008) showed that not all opals are fluorescent. Opal fluorescence can be divided into two classes: blue fluorescence caused by intrinsic oxygen-related defects typical of amorphous silica structures and green fluorescence attributed to uranyl groups (Fritsch *et al.* 2001; 2003), which is believed to be typical of common opals. A low content of U (≥ 1 ppm) automatically induces a green fluorescence response to UV light excitation. This content can reach more than 100 ppm in some deposits (Gaillou *et al.* 2008). Garcia-Guinea *et al.* (2013) measured a uranium amount of 193 ppm in stalactites in Castañar Cave. Thus, detection of opal via UV fluorescence is not systematic but common, as a low content of uranium allows fluorescence emission. However, uranium concentration in Points Cave μ -samples have not been measured due to the particularly thin opal layer and the impossibility of destroying μ -samples.

Table 3. Fluorescence emission peaks for different opal or amorphous silica deposits reviewed in the literature.
Table 3. Pics d'émission de fluorescence pour différents dépôts d'opale ou de silice amorphe examinés dans la littérature.

	Fluorescence emission peaks (nm)				
Othmane <i>et al.</i> 2016	504	524	546	570	-
	504	523	545	573	-
Fritsch <i>et al.</i> 2015	504	524	546	572	604
Brennan & White 2013	502	520	-	-	-
Garcia-Guinea <i>et al.</i> 2013	505	524	543	569	-
<i>This study</i>	501	521	545	572	-

4.4. Opal visual detection under UV light

As explained previously, minerals containing uranyl ions (UO_2^{2+}) have been known to exhibit strong fluorescence marked by specific spectral features and temporal characteristics since at least early 1900 (deNeufville *et al.*, 1981). The bright green fluorescence of uraniferous opal is a well-known characteristic of this mineral phase and has been related to the presence of uranium (Gorobets *et al.*, 1977; Fritsch *et al.*, 2001). To evaluate whether this specificity can be used for opal detection, flake μ -samples were exposed to UV light (see 3.1.1). The results show that μ -samples μ -S-ECA-04 and μ -S-ECA-05 exhibited a bright green-light response (Figure 6). These two sample spectra contained specific features corresponding to the fluorescence emission of uranyl ions. Green illumination was thus associated with the presence of U-opal on these samples. The case of μ -sample μ -S-ECA-03 was more complex. In this case, the absence of uranyl characteristic peaks in fluorescence spectra can be attributed to the low opal occurrence on the μ -sample surface as observed with SEM. However, localized greenish fluorescence could be distinguished on the sample under UV light illumination. This could be explained by a less precise measurement due to a more difficult targeting of opal because of the higher sample size or more scattered opal

distribution. Second, the sample surface was more coloured and had more pictorial matter (as mentioned in §4.1), suggesting decreased ability for detection.

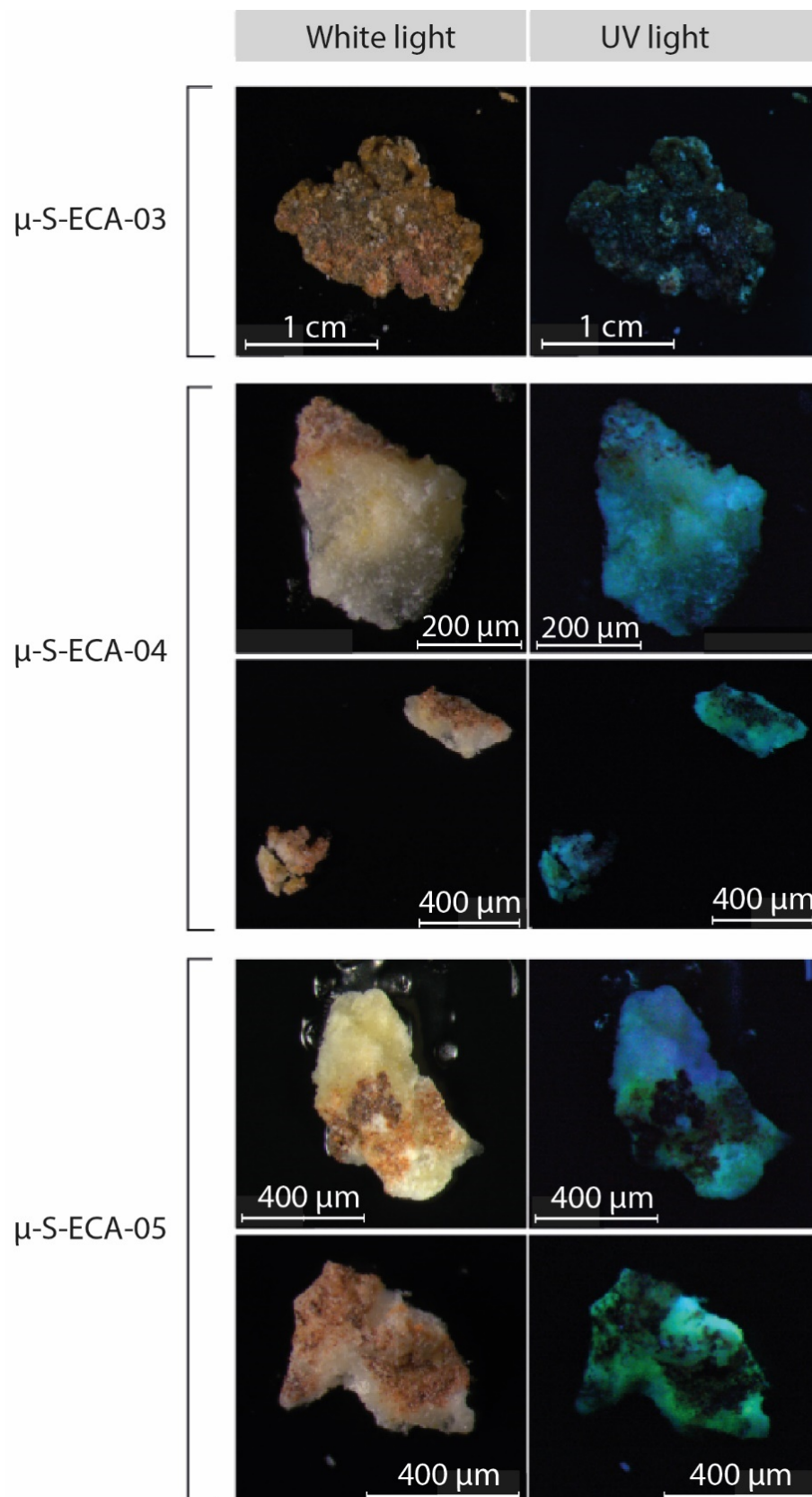


Figure 6. Photographs of μ -samples taken on flakes under white and UV light.

Figure 6. Photographies d'échantillons μ prises sur des paillettes sous lumière blanche et UV.

4.5. From the lab to the cave

4.5.1. Comparison between UV methods

Considering that uranyl green illumination is representative of the presence of opal, detection concordance between spectral and visual techniques was tested at a larger scale by comparing UV-illuminated photographs with UV fluorescence cartography of flake S-ECA-05.

First, to evaluate UV LIF method accuracy at a larger scale, a second experiment was performed directly on flake S-ECA-05. UV fluorescence cartography (100 x 100 μm resolution, Supplementary Information 4) showed a range of spectra presenting uranyl spectral characteristics. Indeed, uranyl peaks present different intensities or ratios compared to the fluorescence signal of the matrix, or cannot be detected. Assuming that the fluorescence signal can be interpreted as a mixing of a uranyl signal with a global matrix signal, opal information was extracted from fluorescence cartography using a mixing algorithm, MCR-ALS, with the MCR-ALS toolbox in MATLAB software (Jaumot *et al.* 2015). MCR-ALS has become a popular chemometric method in solving mixture analytical models. It is based on an additive bilinear model of pure contributions that depends on the concentration and the specific spectral sensitivity of each component. This algorithm can also be applied to obtain quantitative information (Jaumot *et al.*, 2005; de Juan *et al.*, 2014; Zhang & Tauler, 2013).

A singular value decomposition (SVD) method was first applied on the entire raw dataset in the MCR calculation to define the number of initial loadings (Supplementary Information 5). Based on the eigen values obtained with the SVD method, three components were graphically determined as mainly contributors to the fluorescence signal and represent 76.8% of the explained variance. These three initial loadings were then calculated in the MCR-ALS method using the PURE algorithm, a commonly used method to find the purest variable. MCR-ALS was then performed based on these 3 initial loadings on the same entire dataset. Non-negativity constraint was applied for model optimization. For each spectrum, model results ($r^2=99.8$) provided a proportion of each recalculated loading (Figure 7). The second loading represents the uranyl spectrum from which is calculated a proportion of uranyl contribution to the total fluorescence signal. This contribution can be associated with a proportion of opal at the sample surface for each pixel as presented in Figure 7. The other loadings correspond to the contribution of different fluorescence parts in the spectra related to the matrix fluorescence (loading 3) and to some specific highly intense signals at the sample surface (loading 1), probably due to particular minerals.

Evaluation of opal proportions on the sample using this method is, however, subject to certain biases. Similar to the majority of spectroscopic techniques, this detection method can only detect surface deposits. Thus, opal mineralization located under other mineral or organic deposits cannot be detected. Moreover, for quantitative measurement, the measurement sensitivity to changes in sample surface microtopography must be taken into account. As the laser beam is focused on the surface, variations in fluorescence intensities measured on the sample, and thus opal estimations, can be biased by millimetric changes in surface relief.

This method remains effective in extracting information on opal mineralization even on heterogeneous matrices, such as the Points Cave μ -samples. Due to its specific spectral shape, uranyl signal can be easily extracted from mixed fluorescence signals. Mixing algorithms could provide quantitative information if combined with calibration methods. Indeed, the MCR-ALS algorithm has been used for the quantification of different target molecules based on various spectroscopic data (Araya *et al.* 2017; Castro *et al.* 2021; Kumar *et al.* 2018; Mas *et al.* 2010). Regarding sample complexity, calibration protocols based on standard references and prepared or artificial mixtures are difficult to apply (Araya *et al.* 2017). Quantification strategies need to be developed for archaeological materials, reaching a balance between

sample destruction and model result robustness, such as those developed for hyperspectral data. Then, the estimated concentration accuracy will depend on the legitimacy of the assumption made in the quantification strategy (Araya *et al.* 2017).

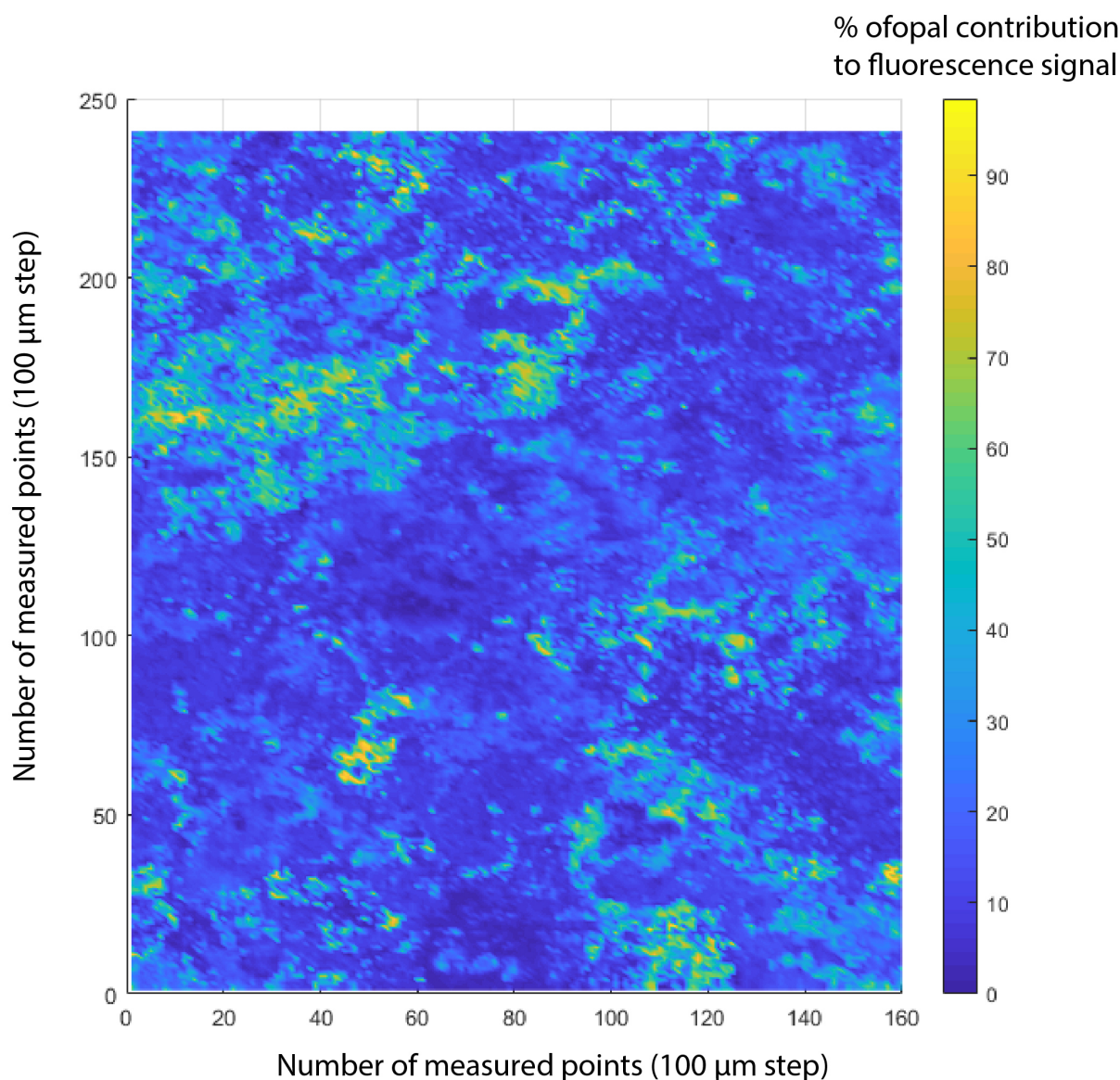


Figure 7. MCR-ALS results: A. Representation of the proportion of opal in the total fluorescence signal calculated from loading 2 on the S-ECA-05 surface, B. Loadings obtained after model optimization.
 Figure 7. Résultats du MCR-ALS: A. Représentation de la proportion d'opale dans le signal de fluorescence total calculé à partir de la charge 2 sur la surface S-ECA-05, B. Charges obtenues après optimisation du modèle.

For field purpose, opal information provided by UV illumination and UV LIF methods were compared. The green component of the RGB image was extracted from UV-light photos. The grayscale image was subtracted to avoid the luminosity effect, and a threshold was applied to the image obtained to select only pixels containing green colour (Supplementary Information 5). To help compare the opal information provided by both techniques, images were aligned using point control selection and geometric transformation functions in MATLAB.

The results show a high correspondence between spectral and visual detection methods (Figure 8). To evaluate the efficiency of the optical method, opal cartographies with different detection limits were simulated by modifying lower boundaries to select pixel associated with

opal presence. These boundaries are based on % of opal contribution to the fluorescence signal for spectroscopic measurements. The best correlation coefficient ($R^2=0.50$) is obtained for an opal contribution threshold of 25% of the total fluorescence signal. Graph D on figure 8 shows that optical method was the most efficient when opal fluorescence emission contributed at least from 20% to 43% of the total fluorescence signal of the sample. If the opal proportion limit for pixel selection is settled too low, the number of pixels selected is too important to provide a good correlation and conversely. For example, 80% of pixels in the image were selected with a limit settled from 8% of opal contribution, providing a correlation coefficient of only 0.24. The rather low correlation coefficient between both images could mainly be explained by the point control selection and geometric transformation applied to readjust them together. Angle disposition of the sample and detector combined with image resolution could have led to difficulties in adjusting the images. Further research in the computational image domain could improve the methodology and enhance the correlation between the two techniques.

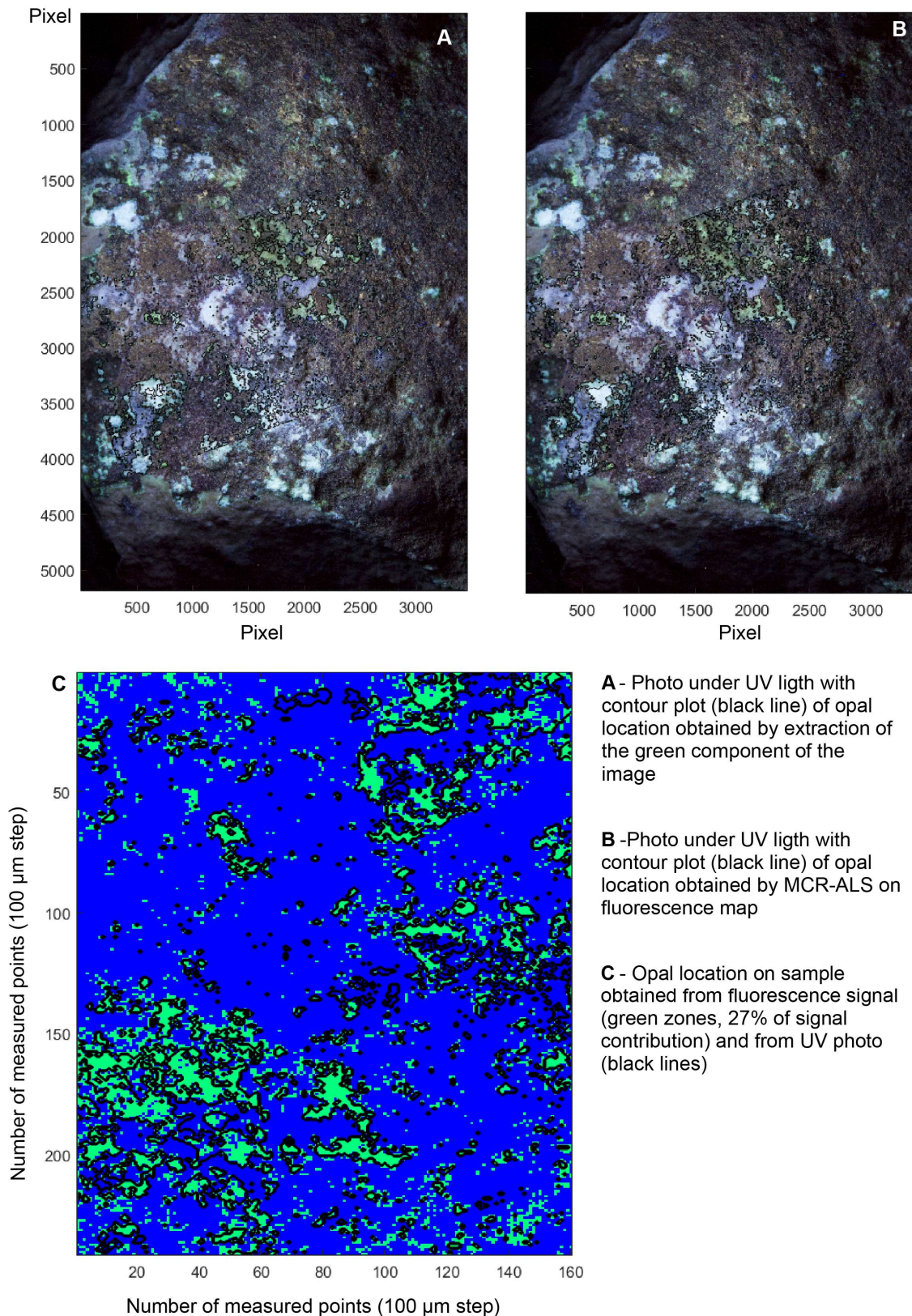


Figure 8. Comparison of detection of opal mineralization on flake S-ECA-05 with UV illumination and UV LIF. A and B. Photo under UV light and contour plot (black line) of opal presence calculated from the green component of the image (at least 8% of green) and from the MCR results (contribution of at least 25% of the total fluorescence) respectively for A and B, after geometric transformation (white square). C. Cartography of pixels associated with the presence of opal comparing A and B (no geometric transformation). D. Correlation coefficient between opal presence images obtained with the two methods (% of opal contribution correspond to lower boundaries applied for pixels selection).

Figure 8. Comparaison de la détection de la minéralisation opale sur l'écaïlle S-ECA-05 avec l'illumination UV et le LIF UV. A et B. Photo sous lumière UV et courbe de niveau (ligne noire) de la présence d'opale calculée à partir de la composante verte de l'image (au moins 8% de vert) et des résultats du MCR (contribution d'au moins 25% de la fluorescence totale) respectivement pour A et B, après transformation géométrique (carré blanc). C. Cartographie des pixels associés à la présence d'opale en comparant A et B (sans transformation géométrique). D. Coefficient de corrélation entre les images de présence d'opale obtenues avec les deux méthodes (les % de contribution de l'opale correspondent aux limites inférieures appliquées pour la sélection des pixels).

4.5.2. Visual detection under UV light: field experiments

An *in situ* photographic protocol that alternated white light and UV illumination was applied to the Points Cave walls. As shown in Figure 1, three rock art panels and one wall devoid of any rock art were investigated. Pictures were taken at both the wall and macroscopic scales.

At the wall scale, the rock art panels investigated (Points Recess, Bi-lobed Signs and Ibexes) displayed large green fluorescent zones when illuminated by UV LEDs (Figure 9). This green fluorescence response was consistent with preliminary results obtained in the laboratory on cave wall flakes and μ -samples from these zones. Thus, UV illumination successfully enabled the detection of opal coatings on these walls, according to the green response observed in photographs.

In contrast, no opal signal appeared clearly distinguishable in photographs of the wall situated a few meters before the art panel (Supplementary Information 7). These photographs displayed greenish fluorescence mixed with other signals, making it difficult to identify and extract green colour with accuracy. Spectroscopic measurements need to be performed to precisely identify the presence of opal.

Macroscopic-scale observations under UV light allowed the targeting of specific deposits or zones on cave walls. In the particular case of the Bi-lobed Signs panel, macroscopic investigation targeting flaked zones revealed a contrast between the well-preserved rock art panels and the afterwards flaked areas. Indeed, the painted surface of the UV photograph (left part of photograph, Figure 9) displayed numerous green fluorescent spots, whereas they were absent from the right part of the photograph. Efflorescence concretions on cave walls, which are widely spread in Points Cave, were also investigated, but the presence of opal has not yet been validated.

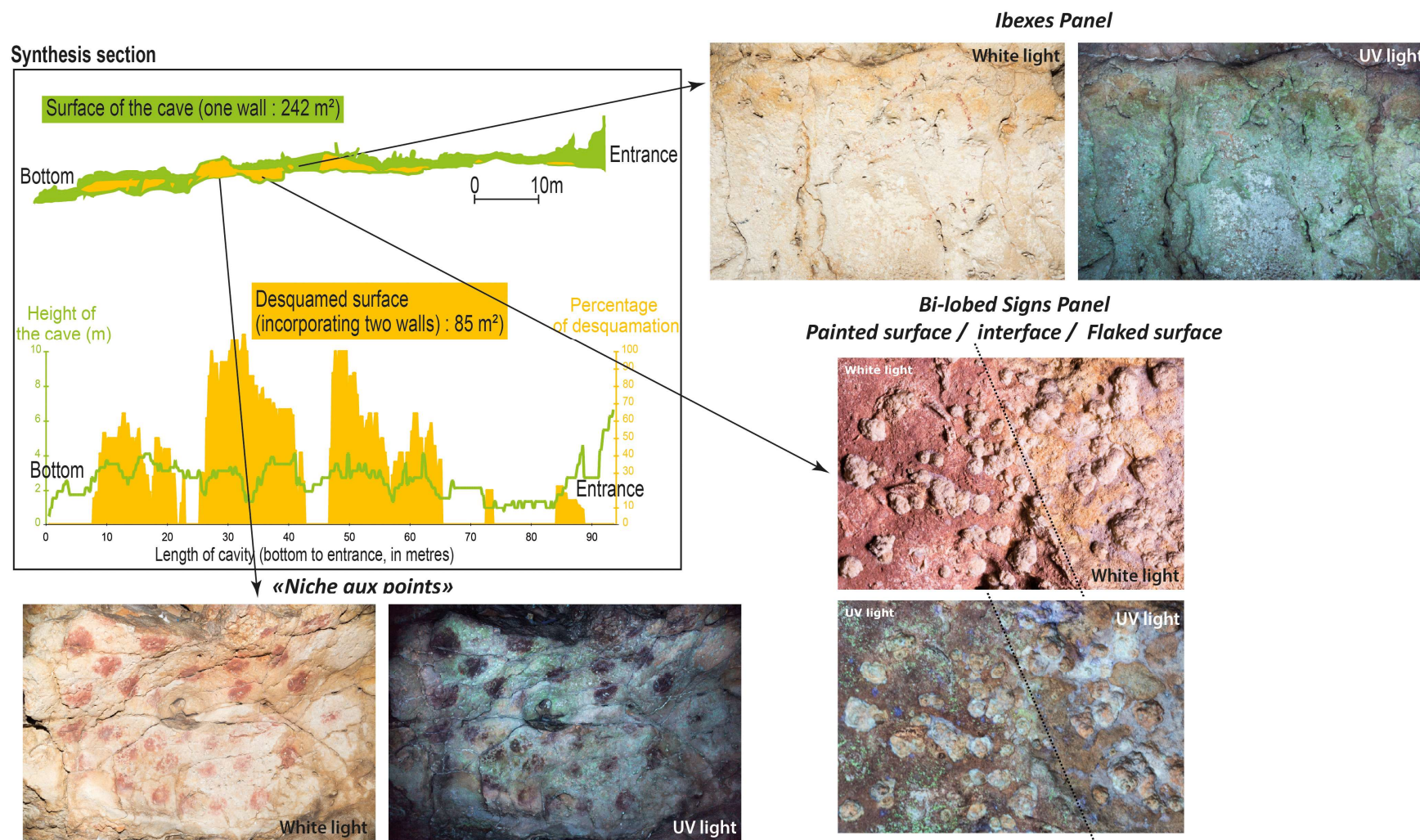


Figure 9. Topographic section of Points Cave with estimated flaking rate of the cave walls according to the distance to the entrance and location of white and UV light photographs according to Jaillet & Monney 2018.

Figure 9. Coupe topographique de la grotte des Points avec estimation du taux d'écaillage des parois de la grotte en fonction de la distance à l'entrée et de l'emplacement des photographies en lumière blanche et UV d'après Jaillet & Monney 2018.

5. Discussion: Implications of opal detection for rock art studies at Points Cave

5.1. Contribution of UV fluorescence techniques

According to the first results obtained using the *in situ* UV illumination method and the preliminary results obtained in the laboratory on cave wall flakes and μ -samples from rock art panels, *in situ* measurements successfully detected the presence of opal in Points cave. Indeed, the green fluorescence observed on Points Recess, Bi-lobed Signs and Ibexes panels suggests that opal development covers a large area of cave walls, especially in rock art sector. Macroscopic observations also help to provide initial insights into rock art-opal interactions, as UV light allows for the targeting of specific deposits or zones on cave walls. It provides complementary information on opal distribution and interaction with archaeological material. For example, investigation of the flaking zone suggests a current absence of extended opal film or opal mineralization, probably due to desquamation occurring on this wall.

However, the opal signal is not always clearly distinguishable, as observed on the wall outside of the rock art sector or on coralloid concretions. Yet, these speleothems are known to contain a silica layer enriched with uranium (Barbarand & Nouet 2020). Alternating layers of silica and calcite or the presence of calcite cover could explain the ambiguous signal. First, this highlights the need for *in situ* fluorescence spectroscopic measurements to validate the photographic identification of the opal mineral phase in cases where the green response is not clearly detectable. Second, colour perception varies greatly from one image to another, even within the same site, making it difficult to evaluate the mineral phase distribution. Even if camera and image treatment parameters influencing colour display (white balance, exposure time, filters, *etc.*) can be easily standardized, lighting variations (LED orientation, position, distance from wall, *etc.*) are difficult to homogenize. As this paper presents preliminary results attempting to validate the detection method, no protocol for colour calibration was applied on UV photographs. Further research on the Points Cave site will be subject to the development of a colour calibration procedure.

Finally, UV illumination also highlights various types of deposits, sometimes not clearly detected under white light. After fluorescence analyses in the laboratory or directly *in situ*, this technique could help to detect other mineral or organic phases. Moreover, it is interesting to note that UV illumination emphasizes the presence of pigments due to their iron composition. Indeed, iron ions are known to have a quenching effect on the fluorescence signal, including that of uranyl ions (Backes 2004; Chen *et al.* 2011; Gaillou *et al.* 2008). Thus, pigments appear clearly as no fluorescence zones on UV photographs, which can be an interesting tool for rock art analysis, such as determination of sampling protocol or dermatoglyph analysis.

UV methodology appears to be an efficient non-invasive tool for the *in situ* identification of U-silica mineralization, and of U-opal mineralization in this case. Due to the green fluorescence resulting from UV excitation, it allows rapid *in situ* detection of the opal mineral phase with direct results and very simple and low-cost equipment. As this method has already been applied with success in some caves, rarely calcareous, for the detection of U-silica complex deposits or concretions, as in Castañar Cave (Garcia-Guinea *et al.* 2013), this study differs in two ways. First, to our knowledge, this method has never been applied to opal detection in a rock art context. Second, we propose *in situ* opal identification based on spectral features for result validation using a portable UV LIF instrument. This instrument is currently in development and could not yet be taken into the cave, but preliminary tests applied to flakes in the laboratory confirm its ability to detect opal. Finally, opal mineral characterization can also be achieved with high-spatial resolution at the microscopic scale with a laboratory UV LIF instrument. Spatially high-resolution fluorescence maps can also provide more information on mineral phase repartition, development and interaction with

other organic and mineral phases present on the sample. Thus, the combination of these two methodologies provides a complete solution for the identification and detection of opal mineralization in both sampling and analytical strategies and *in situ* characterization.

5.2. What we know about Points Cave opal

The results obtained with the analyses presented in this paper and with UV methods applied to samples and *in situ* have provided the first information on opal coatings in Points Cave. Indeed, based on SEM analyses of flakes and μ -samples, Points Cave opal can be identified as the amorphous opal type (type A). XRD is usually performed for precise opal type determination (Curtis *et al.* 2019). However, in the case of Points Cave opal, this analysis is difficult to perform due to the thinness of the opal layer and to the presence iron-rich pigments. Analyses at macro- and microscopic scales suggest thin film deposition as a mineralization type, such as silica skins observed at various rock art sites (Green *et al.* 2017; Watchman 1990). This film appears to be deposited under or at the interface with pigment and other crusts. SEM analyses show that the opal structure encapsulates hematite plates on several samples, suggesting strong interactions with pictorial matter (Figure 4). Further studies must be conducted to understand how these two phases interact, but a preliminary hypothesis regarding dissolution and corrosion of hematite plates by opal can be proposed.

The presence of uranyl ions entrapped in the opal structure was verified by UV spectroscopic analyses. High fluorescence intensities of opal at Points Cave suggest significant content of uranium trapped in the crystalline structure. Similar enrichment in uranium has been found in numerous common opals (Amelin & Back 2006). However, at Points Cave, no quantification of uranium content has yet been realized. Although uranium isotopic measurements were realized on speleothems from Points Cave (stalagmites and coralloids), because of their destructive character, none was applied on flakes, or flake and wall μ -samples. According to the literature, as no minerals containing U⁴⁺ are known to fluoresce, uraniumiferous silica precipitation involves oxidizing conditions sufficient to mobilize U⁶⁺ (Zielinski 1982). For example, Garcia-Guinea *et al.* (2013) explained uranium-bearing opal deposition in Castañar Cave by oxidation of host rocks with meteoritic waters. Zielinski (1982) explained that the initial precipitation of silica was as an amorphous silica gel, with which dissolved uranium coprecipitated before the silica gel dehydrated to form opal. The formation of uranyl-silica complexes is favoured by the natural affinity between aqueous uranyl ions and the silica gel surface, which is very sensitive to pH (optimum range = pH 5-5.5) (Garcia-Guinea *et al.* 2013).

The *in situ* approach shows an important spread of this coating on Points Cave walls, especially on rock art panels. Concerning the undecorated walls a few metres away from the rock art sector, the presence of opal needs to be confirmed by spectroscopic measurement and laboratory analyses. Points Cave chronology is constrained by different periods marked by cave wall evolution under climatic and anthropogenic factors (Monney & Jaillet 2019). The presence of opal mineralization throughout its formation and developmental processes could provide insights into cave chronology. A flaking period was identified subsequent to the rock art phase, resulting from mechanical expansion or desquamation, affecting the deepest zones of the cave, and possibly concurrent with gelifraction at the end of MIS2 (Marine Isotope Stage) or beginning of MIS1 (Monney & Jaillet 2019). As described previously, UV photographs indicated the absence of a well-developed opal film in the flaking zone, suggesting that opal formation occurred principally before this flaking period. This could also explain why photographs taken on walls between the cave angle and the rock art sectors displayed less marked fluorescence, as this zone presents high flaking rates (Jaillet & Monney, 2018) (Figure 9).

The chronology of opal deposits over rock paintings and drawings cannot be confirmed at this stage. SEM observations show that opal mineralization colonizes empty spaces above, below, and inside pictorial matter. These observations suggest an opal deposition subsequent to painting realization, although prior mineralization cannot be ruled out.

Thus, ongoing studies are crucial in completing opal coating characterization in Points Cave. This is a tool in understanding cave wall and pigment interactions throughout the factors and processes of opal formation and other mineralization.

5.3. Opal factors and formation

Mineralization origin and formation mechanisms and factors are clues for understanding past climate and cave wall evolution. As this paper presents preliminary results, only hypotheses for opal origin and formation are presented.

Opal, as a hydrated mineral, is associated with fluid circulation. Thus, water is generally involved in precipitation, and silica has to be in solution before precipitation (Chauviré *et al.* 2017). In Points Cave, the presence of silica coating with high uranium content in the limestone context supports the hypothesis of mineralization originating from groundwater. Wall humidification and water physicochemical properties directly influence the coating formation rate. In very wet sites, the silica skin growth rate can reach 0.25 mm per millennia, whereas occasionally wet sites present a mineralization rate on the order of 0.02 mm per millennia (Aubert *et al.* 2004). Silica coatings can only be formed and preserved if low infiltration volumes occur, as higher volumes favour dissolution of soluble compounds and decrease silica precipitation (Aubert *et al.* 2004).

Chauviré *et al.* (2017) explained that even though opal is found in various geological contexts, three main types of formation can be identified: 1) hydrothermal activity, 2) biological precipitation and 3) continental weathering. As no hydrothermal activity was identified at Points Cave, only the latter two types can be involved in opal formation at this site.

Biological formation of opal in caves is less documented than hydrothermal alteration and continental weathering. Various microbial forms and algae have been observed to be associated with opal in caves, such as siliceous algal diatoms (Northup *et al.* 2001), and are often linked to coralloid concretions. In the case of Points Cave, a few microscopic and cave wall μ -sample observations show undetermined structures containing high carbon content, which could be associated with biological activity. Nevertheless, the hypothesis of biological formation of opal cannot be ruled out or confirmed at this stage.

Continental weathering is defined by rock transformation by meteoric water and a precipitation temperature below 50°C, in contrast to hydrothermal alteration (Chauviré *et al.* 2017). Silica anions released by this process precipitate because of fluid supersaturation due to various changes in conditions, such as pH or temperature (Chauviré *et al.* 2017; Devès *et al.* 2012). Thus, supersaturation of silica solutions may be initiated by a drop-in temperature or pH, or an increase in salinity. When a solution is supersaturated in SiO₂, silicic acid could polymerize to form a colloidal suspension from which amorphous silica can precipitate. Polymerization is controlled by temperature (decreasing T° increases the polymerization rate), degree of supersaturation, salinity and mainly pH (maximum polymerization rate around pH 7.5; minimum polymerization rate under pH 3 and above 9) (Devès *et al.* 2012). One of the most efficient pH-driven mechanisms for silica precipitation involves acidification of highly alkaline solutions (Zielinski 1982). Such alkaline conditions are not common, but in the case of calcite-dominant material in sediment, pH could reach this threshold (Karkanis *et al.* 1999). Freezing temperature has also been shown to help supersaturation, polymerization and rapid precipitation of opal, whereas low or moderate temperatures induce slow polymerization

(months or years) (Devès *et al.* 2012). Cryo-segregation is another reported genesis, caused by moisture freezing on cave walls, which concentrates dissolved salts. They precipitate out in the case of supersaturation of the solution (Devès *et al.* 2012). As opal formation probably occurred during the Late Pleistocene possibly up until the Tardiglacial era and because freezing temperatures have been shown to have modified Points Cave wall topography, a temperature decrease represents a realistic factor for opal formation. However, there is currently a lack of evidence to confirm this hypothesis.

To help understand opal formation, the source of silica and uranium forming this mineralisation can also be questioned. Silica can originate from 1) superficial cover, 2) host rock, or 3) volcanic ash in a continental weathering context (Devès *et al.* 2012). Volcanic ash is an interesting hypothesis, as it is assumed to provide both high contents of silica and uranium, and because the Ardèche region had recent volcanic episodes during the Upper Palaeolithic. Indeed, volcanic eruptions in the Bas-Vivarais region were dated between 23.9 ± 8.1 ka and 31.1 ± 3.9 ka (Nomade *et al.* 2016; Sasco *et al.* 2017). The current alluvial plain and the lowest former alluvial level (+8 m) deposits contain much basaltic material partly derived from Bas-Vivarais lava, which were subjected to intense erosion and weathering in the alluvium terraces (Genuite *et al.* 2021). Thus, the location and chronology of the volcanic activity (approximately 45 km away from Points Cave) represent an interesting origin hypothesis for both silica and uranium contents in opal. However, determining the origin of uranyl-enriched silica solutions is difficult with our current information. Another hypothesis for silica solution origin can be supported by the presence of a marl layer within the Urgonian limestone, which constitutes the cave environment (Sadier 2013); or by pebbles originating from former fluvial deposits originated from the Ardèche River and providing silica by infiltration from the overlaying alluvium terrace or by river deposition (Genuite *et al.* 2021; Mocochain *et al.* 2009; Tassy *et al.* 2013).

Thus, understanding the origin and mineralization factors that influence opal formation could provide information on chemical and physical processes occurring on the cave wall surface. According to Green *et al.* (2017), this knowledge is “crucial for targeted sample collection and the application of a range of dating techniques as well as for the development of conservation strategies”.

5.4. Chronology and dating

Red pigments used in rock art are difficult to date precisely (Aubert *et al.*, 2007). Thus, several studies have proposed indirect dating methods using associated mineral deposits interlaying pictorial matter to date or to provide chronological constraints on rock paintings (Aubert *et al.* 2007; 2017; Watchman 1990), such as opal coatings.

Indeed, in the first place, opal coating could help to precise relative chronology thanks to knowledge regarding climatic and environmental factors controlling its mineralization, and to its distribution on walls in comparison to other deposits and archaeological material.

Amorphous silica skins have been used for radiocarbon analyses based on organic remains trapped by mineralization, such as diatoms or algal matter, on different Australian rock art sites (Morwood *et al.* 2010; Watchman 2000). However, as silica coatings may contain various organic materials, each presenting a specific radiocarbon signature, dating could result in a mixture of different ages by incorporation of younger or older material (Aubert *et al.* 2017; Green *et al.* 2017). In addition, the formation processes of these coatings are not fully understood, requiring great caution when using radiocarbon methods (Aubert *et al.* 2017). Using compound-specific carbon analyses could potentially avoid this problem (Aubert *et al.* 2017) but are more difficult to apply to thin deposits, especially in rock art contexts.

Moreover, opal often contains high uranium contents, which could be used for high-precision dating with methods such as $^{230}\text{U}/\text{Th}$ or U/Pb (Oster *et al.* 2017; Zielinski 1980). Because of opal's ability to concentrate uranium from water while rejecting Pb and Th, amorphous silica is an interesting alternative to carbonate minerals (Amelin & Back 2006). Indeed, $^{230}\text{Th}/\text{U}$ and U/Pb methods have been applied to opal and have provided reliable ages (Neymark & Paces 2013). They enable chronological constraints or dating hydrogenic subsurface water flow, pedogenesis, and processes such as ore formation deposits (Neymark & Paces 2013). They also have been applied in the case of paleoclimate reconstruction in silica speleothem studies (Lundberg *et al.* 2010). Depending on the formation processes, opal coating could thus be used for dating purposes.

In rock art research, dating of amorphous silica deposits could be used as an age constraint for rock drawing events, depending on the pigments and mineral phase superimposition. In their study, Aubert *et al.* (2007) performed U-series dating on a 2.5 mm thick calcite coating using the MC-ICPMS technique, allowing high spatial and temporal resolution. Even though only micrograms of samples are needed, authors (Aubert *et al.* 2007; 2017) have suggested that for samples with U concentrations > 1 ppm, sampling could be largely reduced, and LIBS techniques could also be applied *in situ* with a 100-200 μm diameter ablation spot. As opal concentrates more U than calcite, these techniques appear to be possible. However, for some authors, the application of uranium series dating to silica skins appears difficult to achieve (Green *et al.* 2017). Sampling that provides sufficient intact fragments for LA-ICP-MS analysis without damaging rock paintings is one of the main issues regarding this application. The difficulty of performing closed system conditions and replicability tests for evaluating dating reliability has also been highlighted by authors.

Moreover, methods for opal detection could offer supplemental help before sampling for dating. Indeed, precise targeting of uranium-bearing opal enables identification of pure silica phases in mixed samples, detection of high uranium contents or impurity avoidance. In the case of thin layer deposits, such as in coralloids, sampling could decrease dating accuracy when mixing different layers (Devès *et al.* 2012). Tracking the location of opal phases could avoid this issue by spatially constraining sampling.

5.5. Implication of opal mineral characterization for conservation of rock art material

Taphonomy represents a range of transformations affecting archaeological material that distort archaeological records. Thus, rock paintings have undergone a plurality of transformations impacting pigment longevity, colour and identification (Bednarik 1994; Chalmin *et al.* 2018; Defrasne *et al.* 2019; Huntley 2012; Rousaki *et al.* 2018). At Points Cave, the extensive presence of dermatoglyphs on 47 palm prints is exceptional (Achtelik *et al.* 2019). In fact, to our knowledge, it has no equivalent in European Palaeolithic cave art which raises the question of opal impact on pigment conservation.

If pigment weathering can be influenced by their own properties by inducing changes in surface area, albedo, light transmissivity or moisture (Huntley 2012), mineral deposits have been recognized to be important factors in rock art taphonomy (Chalmin *et al.* 2018). In addition to analytical impacts, knowledge of mineral phases is essential because it informs us regarding the physical and chemical impacts on pictorial matter and whether they favor conservation or degradation effects. Mineral phase characterization is thus an important part of conservation strategies, and adapted identification and analytical methods are needed.

The association of silica skins composed of opal and pigments has been frequently observed at rock art sites, mainly in open-air sites on sandstone and quartzite substrates in Australia or Canada (Aubert *et al.* 2004; 2012; Huntley 2012; Watchman, 1990). Some studies have suggested that pigment binding in silica coatings aids rock art visual preservation

(Watchman 1990), providing a resistant layer to chemical weathering. SEM observations on Points Cave flake and wall μ -samples indeed suggest a strong interaction between pigments and opal, as mineralization penetrates pigment deposits. If the quality of Points Cave paintings tends to corroborate this hypothesis, exfoliation impacting silica skins observed at some sites should also be mentioned, as it could cause removal of associated pigments (Aubert *et al.* 2004; Green *et al.* 2017). Thus, exploiting the observed benefits of silica film deposits for conservation strategies has not yet been proven (Green *et al.* 2017). Furthermore, it has also been observed at open-air sites that silica skins could reduce the colour tone of paintings and drawings at some locations (Green *et al.* 2017).

6. Conclusion

The methodological development proposed in this paper was motivated by the presence of opal in cave art context at Points Cave. Its identification *in situ*, in the laboratory, on centimetric objects and on μ -samples questions the possibility of accessing the specificities of the colouring matter applied on the walls (petrography and geochemistry). Therefore, *in situ* identification of silica coating observed in the laboratory was crucial for further studies on pigment characterization.

The results obtained on flakes and flake μ -samples from Points Cave show that UV LIF is an efficient technique to detect and identify uranyl-silica complexes, even on heterogeneous and complex surfaces. Although opal coating detection is limited by the outcropping nature of the deposit, UV LIF spectroscopy offers a rapid and non-invasive tool that can easily be brought to the site and positioned in front of the rock art panels. A photograph of green bright fluorescence emitted by opal was used here as a method for *in situ* detection of this mineral coating. A comparison with the UV LIF method shows a great correspondence between the two methods. The first tests of UV illumination in the cave highlight the need to develop an accurate measurement protocol, especially to homogenize light and colour, and the need to validate the identification using UV LIF spectroscopy. In addition to further methodological development, UV optical technique shows great potential because it successfully detects the presence of opal and its distribution on rock art panels.

The results obtained with our methodology provide insights into disturbances in the classical *in situ* spectroscopic analyses (pXRF) observed at Points Cave (Chanteraud *et al.* 2021). The identification and characterization of opal coating is thus essential because its impact on *in situ* analysis could disturb the detection of iron-oxide spectroscopic signal in case of Raman analysis for example. For this reason, we propose early, on-site observations combined with sampling of surrounding material, such as fallen flakes from cave walls, as an alternative strategy to i) characterize pigment-associated mineral phases, ii) choose the best site-adapted combination of techniques and devices for *in situ* analyses and iii) define laboratory analytical strategies depending on the pictorial matter environment.

Moreover, optical methods with *in situ* visual detection, such as the UV light illumination method proposed in this paper, represent an interesting tool to add to sampling and analytical strategies. Visualization at a larger scale of the presence and distribution of mineral deposits that could interfere with pigment analyses is a great help in locating sampling or *in situ* measurements to avoid interference.

In addition to analytical impacts, the detection and identification of mineral phases can provide valuable information on the pigment environment and human practice chronology. If speleothems are considered an accurate archive for past climate and environment, other mineral deposits could provide informative records on the setting and evolution of archaeological evidence. Applying a specific methodology for their characterization is thus an efficient tool in improving rock art knowledge.

Thanks to opal mineralization detection utilizing UV methodologies on cave walls, a discussion on its formation and associated factors, such as climatic, hydrologic or geomorphologic conditions over time can be started. Thus, the mineral form described as opal can provide elements on cave natural history.

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Data accessibility statement

All data set (images, composition, spectra) and script (matlab) are accessible for consultation and download on figshare under a CC by 4.0 license.

List of supplementary files

Supplementary Information 1 “S.I.1.” Macroscopic pictures and SEM data of flakes μ -samples):

<https://doi.org/10.6084/m9.figshare.16832593.v2>

<https://doi.org/10.6084/m9.figshare.16832557.v3>

<https://doi.org/10.6084/m9.figshare.16832452>

<https://doi.org/10.6084/m9.figshare.16832605>

Supplementary Information 2 (S.I.2. Macroscopic pictures and SEM data of rock art μ -samples):

<https://doi.org/10.6084/m9.figshare.19142243.v1>

<https://doi.org/10.6084/m9.figshare.19142267.v1>

<https://doi.org/10.6084/m9.figshare.19142276.v1>

<https://doi.org/10.6084/m9.figshare.19142300.v1>

<https://doi.org/10.6084/m9.figshare.19142351.v1>

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<https://doi.org/10.6084/m9.figshare.19142366.v1>

<https://doi.org/10.6084/m9.figshare.19142372.v1>

<https://doi.org/10.6084/m9.figshare.19142375.v1>

<https://doi.org/10.6084/m9.figshare.19142381.v1>

<https://doi.org/10.6084/m9.figshare.19142384.v1>

<https://doi.org/10.6084/m9.figshare.19142387.v1>

Supplementary Information 3 (S.I.3. Single UV fluorescence measurement (raw data)):

<https://doi.org/10.6084/m9.figshare.16837180.v1>

Supplementary Information 4 (S.I.4. UV fluorescence cartography measurement (raw data)):

<https://doi.org/10.6084/m9.figshare.16837324.v3>

Supplementary Information 5 (S.I.5. UV fluorescence data processing (Matlab script)):

<https://doi.org/10.6084/m9.figshare.16837405.v2>

Supplementary Information 6 (S.I.6. pXRF result of in situ analysis at the Points Cave):

<https://doi.org/10.6084/m9.figshare.9791405.v1>

<https://doi.org/10.6084/m9.figshare.9791210.v1>

Supplementary Information 7 (S.I.7. In situ UV photographs):

<https://doi.org/10.6084/m9.figshare.19316588.v1>

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Lumière dans la Grotte: Détection de revêtement d'Opale par illumination et spectroscopie UV en contexte d'art pariétal.

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Abstract:

Les dépôts minéraux des grottes, en tant que produits d'altération, peuvent être considérés comme une source d'informations sur les événements naturels et anthropologiques survenus à la surface des parois. Les revêtements de silice ont été étudiés dans différentes études sur l'art pariétal, mais leur rôle dans la conservation ou l'altération des pigments, ou leur interaction avec la matière picturale, ne sont toujours pas clairs. En outre, ces revêtements sont rarement mentionnés dans un contexte calcaire. Ainsi, le développement de films de silice sur les parois ornées de la grotte aux Points (Gard, France) remet en question l'accès analytique aux spécificités de la matière picturale (géochimie et pétrographie) et l'état de conservation de l'art rupestre dans le contexte de l'étude des pigments. Cependant, les techniques classiques de spectroscopie in situ (pXRF, pRaman) ne permettent pas d'identifier ces revêtements, ce qui empêche également la caractérisation des pigments. Dans cette étude, nous proposons d'utiliser une méthode de fluorescence ultra-violette (UV) pour la détection et la reconnaissance des revêtements d'opale, basée sur les spécificités de fluorescence des complexes uranyle-silice composant ces dépôts. Les premiers développements méthodologiques ont été réalisés en laboratoire avec l'étude d'échantillons et de μ -échantillons provenant du site d'art rupestre de la grotte aux Points. L'identification des dépôts d'opale a d'abord été réalisée à l'échelle microscopique à l'aide d'un microscope électronique à balayage à émission de champ (MEB). Ensuite, une identification spectrale par spectroscopie de fluorescence induite par laser UV (LIF) couplée à une illumination UV a été réalisée sur le même matériau. Les pics bien définis observés dans les spectres d'émission de fluorescence à 501, 521, 545 et 572 nm sont dus aux ions uranyle et valident la détection de l'opale et sa correspondance avec la fluorescence verte observée sous lumière UV aux échelles micro et

macroscopiques. Malgré la nature affleurante de l'échantillon qui limite la détection, la spectroscopie UV LIF offre un outil rapide et non invasif qui peut facilement être amené sur site. Les résultats de la comparaison des techniques UV montrent une grande correspondance entre les méthodes de détection spectrale et visuelle, ce qui permettrait de déployer les méthodes UV dans les grottes. Bien que cette première application souligne la nécessité de développer un protocole de mesure précis, les mesures optiques in situ sous illumination UV révèlent avec succès la présence et la distribution de dépôts d'opale, en particulier sur les parois ornées de la grotte aux Points. Les observations sur la présence et la répartition de l'opale fournissent les premières indications sur l'évolution des parois de la grotte aux Points et les contraintes chronologiques liées au développement de la couche d'opale. Par exemple, les premiers résultats suggèrent un développement du dépôt d'opale principalement avant la période d'écaillage survenue à la grotte aux Points avant la fin du MIS2 (stade isotopique marin) ou le début du MIS1. Au vu des fortes interactions avec les pigments, suggérées par les observations multi-échelles des échantillons et des μ -échantillons, l'impact de la présence d'une couche d'opale sur la qualité de conservation de l'art rupestre de la grotte aux Points est questionné. Cette étude souligne l'importance des observations précoces sur le site, combinées à l'échantillonnage des matériaux environnants, pour la caractérisation préliminaire des phases minérales associées au pigment et comme outil d'aide à la décision pour la définition de stratégies analytiques in situ et en laboratoire en fonction de l'environnement pictural. Ainsi, en développant une méthode de caractérisation non destructive spécifique pour les revêtements d'opale, cette étude ouvre une nouvelle approche pour l'étude de la taphonomie des parois ornées et propose d'utiliser cette minéralisation, à la fois comme marqueur de l'histoire naturelle des grottes, et comme indicateur de leur utilisation par des groupes humains anciens.

Mots-clés : revêtement de silice ; uranyle ; fluorescence UV ; détection in situ ; art rupestre