Identification of nanocrystalline goethite in reduced clay formations.

Application to the Callovian-Oxfordian formation of Bure (France)

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ABSTRACT

The Callovian-Oxfordian (COx) clay formation in the Paris Basin (France) has been the target of many studies investigating the feasibility of deep nuclear waste disposal in a reduced clay formation. To determine the mobility of radionuclides in the host rock formation, modeling of the porewater chemistry, particularly iron solute concentrations, is necessary. Notably, this study aims to understand the supersaturation of Fe(III) oxyhydroxides given by models. Fe(III) oxyhydroxides have been identified magnetically in unpreserved Callovian-Oxfordian samples. In this study, a set of magnetic measurements are used to detect the Fe-bearing magnetic minerals present in the COx clay formation. A core sample from the borehole FOR1118, preserved from air since its collection, is the target of this study. The magnetic measurements performed show that magnetite and goethite are the main magnetic minerals (< 0.2%), together with probable greigite, and occur in low concentrations. Goethite occurs as nanoparticles dispersed in the clayey matrix, and not enclosed in other minerals or in organic matter. It is unlikely that the goethite is an alteration by-product, as particular care was undertaken. This finding resolves the discrepancies between observations and previous modeling results.

Keywords: goethite, nanoparticles, Callovian-Oxfordian clay formation, Paris Basin
Introduction

The Underground Research Laboratory (URL) built in the Callovian-Oxfordian Clay Formation (COx Clay) in Bure, France, is an experimental site used by the French National Radioactive Waste Management Agency (Andra) to test the feasibility of deep nuclear disposal in a reduced clay formation. To assess the durability of materials (e.g. glass, stainless steel, concrete, engineered clay barrier), and to determine the speciation and related mobility of the radionuclides in the host rock formation, it is necessary to model the chemistry of the clay porewater (e.g. Gaucher et al., 2009). Porewater composition models still have some deficiencies that need to be addressed. One of the most problematic issues is the understanding and modeling of iron (Fe) solute concentrations and the subsequent understanding of redox conditions in the formation. This information is mandatory for robust blind simulation modeling of redox perturbations in reduced clay formations (e.g. migration of redox sensitive radionuclides). For instance, predicted water compositions are supersaturated with respect to (Fe(III)) oxy(hydr)oxides (Gaucher et al., 2009). To date, all of the various microscopic and spectrometric techniques used on very well preserved samples of COx Clay in liquid nitrogen failed to predict the presence of Fe oxy(hydr)oxides (Tournassat et al., 2008; Lerouge et al., 2011). This inconsistency between observations and modeling could result from (1) a concentration of Fe(III) oxy(hydr)oxides which is too low to be detected by the different techniques or (2) a problem with the model.

This study aims to test the first hypothesis using a rock magnetic approach in order to detect the magnetic minerals (e.g. iron oxides, oxyhydroxides) possibly present in the reduced clay-rich formation. The main approach used in this paper is based on low-temperature magnetic measurements (from 10 to 400 K). It is non-destructive and allows the identification of magnetic...
minerals, even if they occur in very low concentrations on the order of part per million per volume (ppmv)–part per billion per volume (ppbv).

**Rock magnetism in Callovian-Oxfordian formation: a brief review**

Diamagnetism is a property of all magnetizable minerals. However, diamagnetism is overshadowed if any other forms of magnetism are present (paramagnetism, ferromagnetism).

When a mineral is subjected to a magnetic field (H), it acquires an induced magnetization, M, that is the sum of the transient and remanent magnetizations contributions. Only the ferromagnetic (sensu lato) minerals can retain a remanence. The magnetic susceptibility, \( \chi = M/H \), measured at room temperature under a weak magnetic field, is expressed as: \( \chi = \chi_{\text{dia}} + \chi_{\text{para}} + \chi_{\text{ferro}} \). In argillaceous rocks, \( \chi_{\text{dia}} \) is \( \sim -14 \) µSI (Hrouda, 1986). \( \chi_{\text{para}} \) is similar to \( \chi_{\text{ferro}} \), depending on the ratio of clays (\( \chi_{\text{para}} \)) and ferromagnetic minerals (\( \chi_{\text{ferro}} \)). In the Callovian-Oxfordian Bure claystones, \( \chi \) is dominated by the paramagnetic contribution (\( \chi_{\text{para}} \)) and a positive correlation exists between the magnetic susceptibility and the amount of clays (Esteban, 2006 and references therein). The magnetic susceptibility values and the saturation remanence at room temperature are generally low (< 500 µSI and < 100 µAm² kg⁻¹, respectively) and indicate that a few ferromagnetic grains are present (e.g. Moreau et al., 2005; Esteban, 2006; Aubourg and Pozzi, 2010; Kars et al., 2012). In order to characterize the ferromagnetic contribution, remanence should be investigated.

Room temperature measurements are the most commonly used because they are faster and cheaper than the other techniques. An isothermal remanent magnetization (IRM) is monitored at room temperature with increasing applied magnetic field (generally up to 1-2 T) to achieve saturation. IRM acquisition and hysteresis loops indicate the minerals coercivities, but do not
allow discrimination between minerals with similar coercivity. Nevertheless, statistical analysis and modeling help to discriminate and quantify different IRM components (e.g. Heslop et al., 2002; Egli, 2004). Another way to distinguish mineral coercivities is to impart an IRM with increasing magnetic field values on the three axes, X, Y and Z. This will distinguish between low, medium and high coercivities, respectively (Lowrie, 1990). Then, IRM is thermally demagnetized up to 700°C (maximum temperature generally used). Recognition of the spectrum of unblocking temperature \(T_{ub}\) on a given axis will identify the magnetic mineral. Globally, magnetite is a low to medium coercive mineral \(T_{ub} \leq 580^\circ C\), iron sulfides (greigite, pyrrhotite) are medium to high coercive minerals \(T_{ub} \leq 320^\circ C\) for pyrrhotite), and hematite \(T_{ub} \leq 670^\circ C\) and goethite \(T_{ub} \leq 120^\circ C\) are high coercive minerals. However, this presents some limitations. A recent study by Rochette et al. (2005) has shown that a very high magnetic field (up to 57 T) is necessary to saturate goethite. These high values cannot be reached by magnetometers classically used in rock magnetism laboratories. In addition, thermal demagnetization leads to chemical alteration of the magnetic minerals as a result of heating.

Finally, low-temperature (LT) magnetic measurements (from 300 down to 5 K) appear to be the best method, as they present many advantages. First, because the measurement is performed below room temperature, no sample alteration occurs. Second, similar to the thermal demagnetization, the magnetic minerals are recognized by their magnetic behavior below room temperature. Magnetite shows a magnetic transition at \(\sim 120\) K, called the Verwey transition, which corresponds to a change in symmetry from cubic to monoclinic (e.g. Muxworthy and McClelland, 2000; Özdemir et al., 2002). Hematite shows a Morin transition at \(\sim 250\) K corresponding to a spin reorientation in the C-axis (e.g. Özdemir et al., 2008). Goethite displays a remanence increase from room temperature during cooling, probably as a result of vacancy rearrangement (Dekkers, 1989). Maghemite also shows a remanence increase from room
temperature during cooling, but a smaller increase than that of goethite (generally 10%; Özdemir and Dunlop, 2010). Pyrrhotite shows a magnetic transition at \( \sim 32-35 \) K called the Besnus transition which probably corresponds to a change in symmetry from monoclinic to triclinic (Dekkers et al., 1989; Rochette et al., 1990; Wolfers et al., 2011). This temperature range also corresponds to the magnetic state transition from paramagnetism to antiferromagnetism of Fe-Mn carbonates. Siderite (FeCO\(_3\)) and rhodochrosite (MnCO\(_3\)) transitions occurs at \( \sim 38 \) K and \( \sim 32 \) K, respectively (Housen et al., 1996; Frederichs et al., 2003; Kosterov et al., 2006). The third advantage is that these low-temperature magnetic measurements allow detection of magnetic minerals at very low concentrations (\( \sim 10^{-6} \) g g\(^{-1}\) – ppm to \( \sim 10^{-9} \) g g\(^{-1}\) – ppb).

Recent magnetic studies have been performed on COx claystones from different Andra boreholes in the Paris Basin (Esteban, 2006; Esteban et al., 2006; Aubourg and Pozzi, 2010; Blaise et al., 2013). Esteban et al. (2006) showed that the magnetic susceptibility increases downhole depending on the lithology. The summit of the Callovian-Oxfordian (COx) formation is of lower susceptibility because it is more clay-depleted. The maximum clay zone, located at the middle of the Callovian-Oxfordian formation presents the highest susceptibility values, and rather low remanence. This is due to a high concentration of iron-bearing clay minerals, and low concentrations of ferromagnetic minerals. The base of the COx formation presents high low field magnetic susceptibility values and displays the highest remanence. In these boreholes, occurrences of iron oxides [(titano)-maghemite or magnetite] and iron sulfides (pyrrhotite, greigite, pyrite) have been documented (Esteban et al., 2006). Aubourg and Pozzi (2010) also showed the presence of goethite (probably nanometer-sized) based on LT magnetic measurements on samples from EST211 borehole. This 2010 study is, to the best of the authors’ knowledge, the first to show the occurrence of Fe(III)-bearing oxy-hydroxides in the COx clay formation. The origin of goethite was, however, not discussed. It remains plausible that this
mineral is present due to oxidation through contact with the atmosphere, as no particular care was taken for sample preservation. This present study differs from that of Aubourg and Pozzi (2010) by the analyses of very well preserved COx claystones. The oxidation effect was also tested.

**Materials**

In this study, Callovian-Oxfordian claystones from the Paris Basin (France) are analyzed (Table 1). An additional goethite (α-FeOOH) sample from a laterite of New Caledonia was also measured in order to compare its magnetic behavior to the bulk magnetic signature of COx claystones. The selected goethite was separated by handpicking under binocular.

**Callovian-Oxfordian Claystones**

**General mineralogy of the claystones.** Callovian-Oxfordian (COx) claystones from the Underground Research Laboratory (URL) site in Bure (France) (Gaucher et al., 2004; Andra, 2005; Clauer et al., 2007; Lerouge et al., 2011) were extensively studied for their chemistry and mineralogy. COx Clay had undergone limited burial (600 m) and diagenetic (T < 50°C) processes (Pellenard et al., 1999; Elion et al., 2005; Clauer et al., 2007). Its mineralogy consists of a dominant clay fraction (> 40 %) associated with carbonates (< 25 %) and quartz (< 25 %), with minor K-feldspars (< 2 %) (Tournassat et al., 2007; Table 2). Clay minerals are dominantly illite-smectite mixed-layer minerals, illite, chlorite and kaolinite with minor glauconite, muscovite and biotite (Pellenard et al., 1999; Gaucher et al., 2004; Pellenard and Deconinck, 2006). A maximum clay zone (MCZ) is defined at 486-489 m depth in the borehole EST-205. Beneath and within this
MCZ, the clay fraction is high (40-60 %) and illite-smectite mixed-layer minerals are ordered following a Rechweite parameter (junction probability) of 1 (ISR1). Illite-smectite mixed-layer minerals are dominantly illitic in the whole formation, and in addition to the disappearance of discrete smectite with increasing burial depth, the transition from a random (R0) to a R1 stacking mode may be driven by interaction with organic matter (Claret et al., 2004). Above the MCZ, the clay fraction decreases towards the top in favor of a silty, then a calcareous-rich, fraction (Gaucher et al., 2004; Andra, 2005). Kaolinite and biotite disappear, and the ordered illite-smectite mixed-layer minerals (ISR1) are replaced by randomly interstratified illite-smectite mixed-layer minerals dominated by smectite (ISR0). Chlorite is tri-octahedral, iron-rich, and of the chamosite-type, according to the Hey classification (Hey, 1954; Lerouge et al., 2011).

The carbonate fraction is composed predominantly of calcite, with minor dolomite, ankerite and siderite. Accessory phases (< 2 %) are pyrite, rutile/anatase, celestite, and apatite. Organic matter content is low (< 0.6 %), and immature in the petroleum sense. Amongst all the mineral phases, the main Fe carrier phases are, in order of decreasing iron content, pyrite, glauconite, biotite, chlorite, muscovite, and illite/illite-smectite mixed layers (Tournassat et al., 2008; Lerouge et al., 2011).

**Studied claystones.** Two core samples (EST26479 and K119), deemed to be representative of the COx Clay Formation, were selected for this study. Sample EST26479 is a preserved core sample from the FOR1118 borehole collected at depth -124.06 m (ZNFG; NGF: Nivellement Général de la France) in 2009. The sample was cleaned and conditioned under nitrogen gas in a gas cell on the field immediately after drilling. It was then transferred and kept into a glove box in a nitrogen atmosphere (O₂ concentration lower than or equal to 1 ppm) (modified from...
Gaucher et al., 2009). Sample K119 is an unpreserved core sample of the EST-205 borehole collected at depth -111.50 m (Z_{NGF}) in 2000 and always kept in open atmosphere. Both samples belong to the C2b1 sub-unit and are almost close to the gallery level. Their mineralogical composition is summarized in Table 2.

In order to determine if pyrite and organic matter contain enclosed magnetic minerals (e.g. iron oxides), they were separated from the claystone argillaceous matrix by handpicking. Two samples were considered (Table 1). A pluri-cm–scale pyritized ammonite collected in 2008 during the opening of a new gallery in the Bure URL was selected to obtain a representative fraction of pyrite associated with minor amounts of organic matter. An organic matter-rich fraction was separated by handpicking in the preserved EST26479 sample.

Sample preparation for low temperature measurements

The selected massive goethite, non-preserved from atmosphere, was first crushed into ~1 mm-sized grains and washed three times with 95 % alcohol in an ultrasonic bath (5 minutes). It was then dried at 40 °C, before being measured in this present study.

The preserved EST26479 bulk sample was crushed in two different ways: into coarse granulates and fine-grained powder. Coarse granulates and a first aliquot of fine-grained powder were prepared and inserted in two gelatin capsules in the glove box under nitrogen atmosphere. They were then encapsulated and sealed in a glass vial prior to removing it from the glove box. Glass vials were opened just before the magnetic measurement. Another aliquot of fine-grained powder was prepared in air 24 hours before the measurement in order to test the oxidation effect after 24 hours.
The pyrite sample used for magnetic measurements was obtained by crushing the pluri-cm–scale pyritized ammonite from a gallery opened in 2008 at the Bure URL, and its inner part was selected. The sample was broken into mm-sized grains in air and rapidly wedged inside the gelatin capsule with quartz wool just before the measurement, in order to obtain fresh surfaces and to limit, as much as possible, sample oxidation.

The unpreserved K119 bulk sample and organic matter extracted from sample EST26479 were encapsulated in air just before measurement.

**Methods**

**Raman spectrometry**

Micro-Raman measurements were performed at BRGM with a Renishaw InVIA Reflex microspectrometer coupled to a DMLM Leica microscope equipped with 4 objectives with x5, x20, x50 and x100 magnification. The excitation laser was an argon laser (\(\lambda_0 = 514.5 \text{ nm}\)). Each spectrum was acquired by two signal accumulations with an acquisition time of 10 seconds per analysis, over a spectral range which depends on the mineral studied. Calibration was done using the 520.4 cm\(^{-1}\) line of silicon. Instrument control and Raman measurements were performed with Renishaw WIRE\textsuperscript{TM} software.

**Magnetic measurements**

**Low-temperature measurements.** In order to determine the rock magnetic properties of the samples, low-temperature magnetic measurements were performed using SQUID cryogenic
magnetometers called Magnetic Properties Measurement System (MPMS) at the Institut de Physique du Globe de Paris, France and at the Institute for Rock Magnetism, University of Minnesota, USA. For this study, the evolution of a saturation isothermal remanent magnetization (SIRM) obtained with the application of a 2.5 T magnetic field at room temperature (RT-SIRM at 300 K) was monitored. Though the term “saturation” is used, it should be noted that it is not a true SIRM, as neither all goethite nor hematite will be saturated at 2.5 T. Two measurement sequences, hereafter referred to as A and B, were used and are described below (see also supplementary material).

Sequence A consists of a cooling/warming cycle of a SIRM imparted at room temperature, for which the value at 300 K is named M1, from 300 K to 10 K. The sample was cooled down to 10 K and then warmed to 300 K in zero magnetic field (trapped field < 0.1 µT). This 300-10-300 K sequence was used to determine the nature, and an approximated crystal size, of the magnetic minerals present in the samples.

Sequence B is designed specifically to underline goethite and to quantify its abundance. This sequence is based on Guyodo’s et al. (2006) protocol. Before applying sequence B, the sample was primarily measured with sequence A. Then at 300 K, it was warmed to 395 K, above the Néel temperature of goethite which is ~393 K (120°C; Özdemir and Dunlop, 1996). At this temperature, the remanence carried by goethite is lost, as goethite becomes paramagnetic. At 395 K, the sample was cooled down to room temperature in a 2.5 T magnetic field. It is generally designed to impart magnetization on a wide range of grain size of hard coercive minerals, i.e., maximize the amount of magnetization carried by goethite. At 300 K, the magnetic field was switched off. The remanence at this temperature step is called M2. Then the sample underwent a second cycle of cooling and warming from 300 K to 10 K and from 10 K to 400 K, similar to sequence A. At 400 K, all the remanence carried by goethite was removed. Finally, a 400-10-300...
K cycle ended the sequence. The two last cycles were performed in zero magnetic field. At the end of sequence B, the obtained remanent magnetization is referred to as M3. The difference between M2 and M3 is a proxy of the goethite concentration in the sample, as M2 is believed to concentrate all goethite and M3 is free of goethite (Guyodo et al., 2006). The approach used here differs from the Guyodo’s et al. (2006) protocol by the non application of AF demagnetization at 200 mT after the measurement of M2 to remove the contribution of ferrimagnetic minerals with coercivity lower than 200 mT (e.g. magnetite). The sequence used in this study was designed to enhance the signal carried by goethite without removing the sample from inside the MPMS and to counterbalance the absence of a strong demagnetizer similar to the one used by Guyodo et al. (2006).

Finally, a third sequence was used on a COx sample and involved imparting an IRM at 10 K, called LT-SIRM, with a 2.5 T magnetic field. The sample was then warmed up to room temperature in zero magnetic field.

Magnetic measurements were performed on about 300-700 mg of sample (fragments or powder) wedged inside 0.5 x 1.4 cm gelatin capsules.

**Room and high temperature measurements.** To complete low-temperature magnetic measurements, room and high temperature measurements were also performed on the FOR1118 sample. Several aliquots were used. For the thermal demagnetization experiment, the FOR1118 samples were conditioned in plaster with a cylindrical shape. An IRM was imparted, by using an impulse magnetizer ASC IM-10-30, on the X, Y, Z axes with 0.08 T, 1.2 T and 2.8 T magnetic fields, corresponding to soft, medium and hard components, respectively (Lowrie, 1990). The specimens were then thermally demagnetized from 80°C to 600°C in an oven in the shielded room of the paleomagnetic laboratory of Université de Montpellier 2, in order to determine the
Curie/Néel temperature of the magnetic minerals present. The remanence was measured after each heating step with a 2G SQUID cryogenic magnetometer. One hysteresis loop with a 400 mT saturation field was measured with a Princeton Vibrating Sample Magnetometer at room temperature at the Institute for Rock Magnetism.

Results

Goethite

Goethite was measured with sequence B. The remanence values M1, M2, M3 are reported in Table A2 of the supplementary material. The data show that the goethite remanence acquired at room temperature (M1 ~ 8.10^{-3} Am^2 kg^{-1}) increases about four times as temperature was reduced down to 10 K (Figure 1a). This magnetic behavior, associated with a high percentage increase, is typical of goethite according to the literature (e.g. Dekkers, 1989). When warming back to 300 K, the remanence shows characteristic quasi-reversible behavior. After the 300-395 K heating-cooling phase under a 2.5 T magnetic field (Figure 1a), the imparted SIRM at room temperature (M2 ~ 3.10^{-2} Am^2 kg^{-1}) was about four times higher than M1. The heating-cooling step under the magnetic field is then very useful to mobilize the goethite grains. From 300 to 395 K, the remanence decreases by 95 % when passing through the Néel temperature of goethite (120°C-393 K; Özdemir and Dunlop, 1996). When performing a 400-10-300 K cycle, the remanence shows an almost flat (constant) reversible behavior. All the goethite grains are thus demagnetized. About 15 % of the remanence (M3 ~ 1.10^{-3} Am^2 kg^{-1}) is nevertheless still present, suggesting that one or more magnetic phases with Curie/Néel temperature higher than 395 K occur (e.g. hematite) (Guyodo et al., 2006).
Callovian–Oxfordian claystones from the Paris Basin

**Bulk samples.** The room temperature hysteresis loop shows that the contribution of ferromagnetics (sensu lato) is very small (Figure 2a). Saturation is reached at 400 mT suggesting that the occurrence of low coercive minerals (magnetite or maghemite) (Figure 2b).

This is supported by the thermal demagnetization of IRMs showing that most of the remanence is carried by the X axis (low coercive phase) (Figure 3). The drop in remanence at ~250 °C suggests the presence of iron sulfides (greigite) (Roberts et al., 2011). Remanence is totally removed by 580 °C, indicating that magnetite is present. In specimen 2, a drop of the remanent magnetization is observed at ~120 °C on the soft component (Figure 3). This drop could correspond to the removal of a viscous component. No clear drop is identified on specimen 1 (Figure 3).

To better characterize the magnetic mineralogy of the COx claystones, additional measurements were performed. Preserved fragments of EST26479 are measured at low temperature. The remanence evolution through a cooling-warming cycle in zero magnetic field (sequence A) shows that the remanence increases by ~30 % from 300 to 10 K, suggesting the occurrence of goethite (Dekkers, 1989). The remanence drop of the Verwey transition of magnetite is not recovered when warming back to room temperature (Figure 4a) (Muxworthy and McClelland, 2000; Özdemir et al., 2002).

In order to corroborate this preliminary result, a crushed EST26479 specimen is measured with the sequences A and B. The cooling-warming SIRM cycles before and after the 395 K heating phase under a 2.5 T field are quasi-superimposed (Figures 4b and4c). From 300 to 10 K, the remanence increases by ~30 %, suggesting goethite. When warming back to room temperature, the curves are reversible. After heating the sample to 395 K in zero magnetic field, ~76 % of the EST26479 remanence remains. In other words, the fourth of the remanence is lost when heating
the sample to the Néel temperature of goethite (Özdemir and Dunlop, 1996). The fact that a non-negligible part of the remanence remains at the end of sequence B (after heating) suggests that one or more magnetic phases are present in the EST26479 sample in addition to goethite. These additional magnetic phase(s) have Néel/Curie temperatures higher than 400 K, such as (titano)-magnetite, maghemite or hematite. Neither the Verwey transition of magnetite nor the Morin transition of hematite is, however, observed on low-temperature measurements (Figures 4b and 4c).

Nevertheless, maghemite may be present in a minor amount. Maghemite and goethite are both magnetic minerals that could result from the oxidation of reduced iron-bearing phases. Oxidation is tested by considering two distinct COx samples. One EST26479 specimen left in air during 24 h after crushing is measured at low temperature (Figure 4d). The remanence regularly increases by ~46 % down to 10 K, typical of goethite. The cooling-warming cycle is reversible. The second sample came from the K119 borehole, and was left in air since its collection in 2000 (Figure 4e). The results obtained are similar to the 24h air EST26479 samples. For 24h air EST26479 and K119 claystones, no other magnetic minerals are identified with low-temperature magnetic measurements. Both analyzed samples thus show a ~50 % increase in remanence from 300 to 10 K, typical of goethite. This is higher than the ~30 % increase for the preserved samples (Figures 4a, 4b, 4c).

Pyrite and organic matter separates. Pyrite and organic matter are present in small amounts in the Callovian-Oxfordian clay formation (<1%) (Table 2; Gaucher et al., 2004). However, they represent a significant reduced-iron-rich fraction, which could be rapidly oxidized under oxic conditions to produce goethite. Measuring an SIRM on the pyrite-rich separate from a preserved pyritized ammonite (Figures 5a and 5b) was not possible because of the very low concentration
(or absence) of ferromagnetic minerals capable of retaining a remanence. Micro Raman spectra on pyrite fragments do not allow identification of any iron oxides or hydroxides (Figure 5c).

An organic matter-rich separate (with probable pyrite content) from the EST26479 sample was analyzed (Figures 5d and 5e). No evolution of the SIRM imparted at room temperature is observed when cooling down to 10 K (Figure 6). The warming curve shows irreversible behavior from ~60 K. This could suggest the presence of a magnetic phase. Neither magnetic transitions nor any particular SIRM increase during warming is identified. This means that magnetite, hematite and goethite can be excluded. Then, remaining candidates could be oxidized magnetite or maghemite. Based on the study by Esteban et al. (2006), maghemite is likely, although oxidized magnetite cannot be ruled out in the absence of complementary data.

Discussion

Similar to previous studies (Esteban et al., 2006; Aubourg and Pozzi, 2010; Blaise et al., 2013), magnetite is identified in preserved COx claystones when monitoring artificial remanence at low temperature (Verwey transition at 120 K, i.e. -153 °C) (Figure 4a) and at high temperature ($T_{ub} < 580$ °C) (Figure 3). Some magnetite grains might possibly be newly-formed during diagenesis. In the high temperature experiment, a break-in-slope near 250 °C suggests the presence of iron sulfides (Figure 3). Such behavior was also reported by Esteban (2006). In practice, it could be greigite or pyrrhotite. However, low-temperature experiments did not show evidence for the Besnus transition of pyrrhotite (at ~32 K) (Figures 4a, 4b). Thus, the iron sulfide is probably greigite. This mineral is a classic product of early diagenesis (Roberts et al., 2011). It is worth noting that the magnetic assemblage of both magnetite and greigite is expected to be representative of immature claystones according to Aubourg et al. (2012).
The low-temperature magnetic analysis revealed the diagnostic signature of goethite contribution with an increase of remanence from 30% in preserved claystones to 50% in claystones unpreserved from the atmosphere during cooling (Figure 4). It remains possible that a portion of this increase in remanence from 300 K to 10 K is due to the contribution from maghemite. In any case, goethite is observed in preserved COx claystones. Similar goethite signatures have been reported by Aubourg and Pozzi (2010) and Blaise et al. (2013). By contrast, high-temperature analysis failed in the firm identification of goethite in preserved COx claystones. A slight drop at 120°C is observed when demagnetizing the soft component of IRM (Figure 3) but it is unlikely that this is due to goethite because this mineral has a hard coercivity (Rochette et al., 2005). Therefore, this present study confirms that high-temperature techniques in rock magnetism are not adapted to detect goethite contribution, but low-temperature (< 300 K) measurements appear very useful.

Other magnetic phases are, however, probably present, but have not been clearly identified by magnetic measurements. Maghemite as an accessory phase could not be dismissed, as pyrite and organic matter extracts may show the occurrence of maghemite (Figure 6). The theoretically well-marked Morin transition of hematite (~250 K; Özdemir et al., 2008) is not observed in the studied samples. In addition, the high temperature measurements performed in this study do not allow the recognition of hematite (Figure 3).

The different magnetic analyses conducted in this study provide evidence that goethite, magnetite and possibly greigite are the main magnetic minerals in the COx samples which were carefully preserved from any oxidation effects.
Contribution of goethite in claystones

The relative concentration, $c$, of goethite (with size above the blocking volume, $V_b$) in COx claystones could be approximated by $c \sim (M_2-M_3)/M_{rs}$ with $M_{rs}$ representing the saturation remanent magnetization of goethite ($M_{rs} = 0.05 \text{ Am}^2 \text{ kg}^{-1}$; e.g. Dunlop and Özdemir, 1997). $M_2$ is the IRM value at 300 K after the 2.5 T cooling phase between 400 and 300 K. $M_3$ is the IRM value at 300 K after the 300-400 K heating/cooling phase in a null magnetic field. In this equation, only goethite is assumed to contribute to the measured SIRM. For the EST26479 COx sample (Figure 4b), $M_2$ is $1.58 \times 10^{-4} \text{ Am}^2 \text{ kg}^{-1}$ and $M_3$ is $1.21 \times 10^{-4} \text{ Am}^2 \text{ kg}^{-1}$ (Table 3). The estimated goethite concentration is then $< 0.1 \%$ of the mass. Similarly, a simplified calculation of the goethite concentration, $c$, might be used for samples which were measured only with sequence A. This is expressed as $c \sim M_1/M_{rs}$. For the preserved EST26479 fragments (Figure 4a), the estimated concentration is $< 0.2 \%$ (similar to the K119 sample). Magnetite and other magnetic minerals might also occur. The $M_{rs}$ of soft magnetite is $\sim 10 \text{ Am}^2 \text{ kg}^{-1}$ (Maher and Thompson, 1999), i.e., 200 times higher than that of goethite, leading to an overestimation of the goethite (above $V_b$) concentration. Such low concentration values explain why goethite has not been identified before in the routine mineralogical analyses at room temperature. For instance, X-ray diffraction has a threshold value of about 1 $\%$ when the mineralogical assemblage is complex, as it is the case in clay-rich rocks, and detection is further complicated when phases are nanocrystalline, because the intensity diffracted at each lattice point is distributed over a large angular range.

However, the concentration of goethite occurring in sizes below $V_b$ (superparamagnetic grains) may be higher. The monitoring of the LT-SIRM of a preserved EST26479 sample displays a significant decrease of $\sim 60 \%$ from 10 to 50 K (Figure 7). This suggests the presence of small
particles that unblock very rapidly (i.e. nanoparticles). These small particles could be magnetite, goethite or other nano-magnetic minerals (greigite, maghemite). In addition to their low concentrations in COx samples, the magnetic minerals occur in small sizes. Both features together make them hardly detectable.

Finally, magnetic measurements carried out on both pyrite and organic matter extracts have shown that they carry a small remanence. Iron oxides such as magnetite and maghemite, and iron sulfides (greigite) can be evoked to explain the magnetic behavior observed in Figure 6. Based on the available data, none of these can be ruled out. Magnetite and/or maghemite are more likely as an alteration by-product because pyrite oxidizes into iron oxides. These iron oxides can also be present as impurities resulting from the separation protocol, or impurities enclosed in pyrite and organic matter. The measurements performed in this study do not permit the distinction between these different hypotheses, and their origin is still debated. No goethite is identified in these two extracts, suggesting that goethite is dispersed within the clayey matrix, rather than being locally enriched as impurities or being consecutive oxidation by-products.

**Implications for the modeling of pore water chemistry in reduced clay formations**

The finding of small amounts of goethite and magnetite in COx samples preserved from atmosphere are important for our understanding of the redox conditions which can be found in the pore water of clay-rocks. Amongst pore water chemistry parameters, redox potential is one of the most important because predictions of the solubility and the mobility of redox sensitive species (e.g. uranium) rely on the accuracy of its determination. To date, it is difficult, if not impossible, to measure directly the redox potential of the pore water in a clay-rocks sample. The
measurement of the redox potential of seepage water flowing from the clay formation to an equipped borehole is subject to experimental artifacts: biological activity, including sulfate reducing bacteria metabolism, has been always detected in such conditions (Vinsot et al., 2008).

This biological activity is not expected in-situ where the pore space is too limited for bacteria to be active. Consequently, modeling the redox potential of the pore water is the only way to estimate it, and the presence of goethite and magnetite in the mineralogical assemblage is an important modeling constraint.

A comprehensive description of the modeling procedure that has been applied to the Callovian-Oxfordian formation can be found elsewhere (Gaucher et al., 2009). The modeling procedure is based on the finding of chemical controls that constrain the system at equilibrium following Gibbs’ Phase Rule. In pore water modeling, the concentrations of the solid phases are not important, only the presence (or absence) of a phase at equilibrium provide useful constraints.

Sulfate is present in the pore water of Callovian-Oxfordian claystone and its concentration is well constrained by measurements (Vinsot et al., 2008). Its concentration in solution is also in agreement with equilibrium with celestite (SrSO₄). Redox potential can be calculated by considering the S(VI)/S(-II) redox couple, provided that the concentration of S(-II) can be estimated. Unfortunately, the S(-II) concentration cannot be measured and, if so, it is likely to be subject to bacterial activity perturbation. If equilibrium with pyrite, that is in contact with the pore water, is assumed, it is possible to estimate the S(VI)/S(-II) redox balance. Again, this is only possible if the Fe(II) concentration is known. This is achieved by assuming equilibrium of the pore water with siderite (FeCO₃), all other pore water parameters being controlled by other constraints (e.g. equilibrium with calcite, quartz etc.). Applying this modeling procedure has led to pore water compositions that are over-saturated with respect to goethite and other Fe(III) (hydr-) oxides (Gaucher et al., 2009) where these phases had not been observed until now on
well-preserved samples, using, for instance, XRD and 77 K Mössbauer spectrometry (Tournassat et al., 2008). So far, two main explanations have been put forward to explain this inconsistency: (i) an inaccuracy in the Fe phases’ solubility products as tabulated in the used database and (ii) an inaccuracy originating from the deviation of the composition of the actual iron carbonate phase from that of the pure siderite considered in the modeling approach (Gaucher et al., 2009, Pearson et al., 2011). A third explanation could be that pyrite and sulfate are not at equilibrium. Sulfate species are known not to be reduced easily at low temperatures through abiotic processes (Truche et al., 2010). This study brings a part of the answer. Magnetite and especially goethite (Fe(III)-bearing mineral) have been identified in the COx samples. Following this finding, a new pore water modeling has been conducted (Table 3) using the same mineral solubility database as the one used by, and available in, Gaucher et al. (2009).

Table 3 shows that considering different combinations of Fe/redox controlling phases has little effect on the modeled pH values which span a range of values from 7 to 7.5. The redox potential values (expressed here as pe = 0.059 Eh) are not much affected by the different hypotheses, with the exception of the goethite/siderite and goethite/magnetite calculation cases. For the three other systems, pH/pe couple values correspond to a single O₂ or H₂ fugacity value (an increase of one pH unit leads to a decrease of one pe unit). Table 3 shows also that all Fe phases identified in the formation cannot be at equilibrium with the pore water but this is a logical result of the direct application of a model following Gibbs’ Phase Rule: the number of controlling phases in the system cannot exceed the number of degrees of freedoms in those calculations (see Pearson et al., 2011). Even so, the finding of two phases at near-equilibrium could indicate that these two phases can co-exist from a thermodynamics point of view. Here, it is apparently not the case: the co-existence of goethite and magnetite at equilibrium would lead to pyrite oversaturation but also to a Fe solute concentration that is not in agreement with the range of values measured in the
seepage waters from the formation (15 ± 11 µmol L⁻¹ according to Gaucher et al., 2009). This result could indicate that one of the minerals goethite or magnetite is metastable in the investigated samples, or that the considered solubility products are not sufficiently accurate to draw any definitive conclusion. Indeed, the present study shows that goethite must be a nanocrystalline phase in the investigated samples. If so, its solubility might deviate from the value tabulated in databases that are representative of well-crystallized minerals. The solubility of goethite in our sample might therefore be larger than the tabulated value in the database (Electronic Appendix 1 in Gaucher et al. 2009). The line in Table 3 corresponding to the system magnetite/pyrite shows that a slight change in goethite solubility could be sufficient to achieve equilibrium between magnetite/goethite and pyrite. Moreover, for this calculation, the Fe concentration is consistent with the range of values measured in seepage water (Gaucher et al., 2009) and the siderite saturation index is only slightly negative (Table 3). This siderite undersaturation can be linked to the exact nature of the Fe carbonate: a sideroplesite whose solubility might be lower than the solubility of pure siderite (Tournassat et al. 2008, Lerouge et al., 2013).

It is thus possible to reconcile the observations made in this study (i.e. the presence of nanocrystalline goethite and magnetite in pristine samples) with the pore water modeling results. A window of pH/redox conditions ranging from pH 7 to 7.5 and pe from -3 to -3.9 can be also given for the Fe(II)/Fe(III) redox controlling system.

Conclusive Remarks

Porewater modeling in Callovian-Oxfordian claystones from the Paris basin has predicted the occurrence of goethite; however, to date, standard observations on fresh or well-preserved samples have failed to detect it. In this study, the low-temperature magnetic properties of
Isothermal remanent magnetization are used to detect trace amounts of goethite. Nanoparticles of goethite, in addition to magnetite and possibly greigite, are detected in both preserved and unpreserved claystones. This strengthens the idea that goethite is naturally present in COx claystones and is not the result of oxidation in contact with atmosphere. This goethite is not detected in pyrite and organic matter extracts. Goethite is probably dispersed within the clay fraction. The low-temperature magnetic technique appears to be appropriate to detect nanoparticles of goethite in sediments.

**Acknowledgments**

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REFERENCES


FIGURES CAPTIONS

Figure 1: Evolution of the RT-SIRM for the reference goethite during three successive cooling-warming cycles. (b) is a zoom of (a) in the 200-360 K temperature range. The increase in remanence from 300 to 10 K is 75 and 73% for cycles 1 and 2. A slight Morin transition of hematite is observed at ~250 K. Note that the remanence is almost entirely removed after the last heating step at 395 K (cycle 3).

Figure 2: a) Uncorrected and b) corrected hysteresis loop at room temperature for bulk Callovian-Oxfordian claystone from FOR1118. Note the high paramagnetic contribution in the sample. Hysteresis loop was corrected by assuming saturation above 70% of Hmax. Saturation is reached at 400 mT.

Figure 3: Thermal demagnetization of IRM showing soft, medium and hard components. Note that more than 50% of the initial remanence is lost by 250°C suggesting the presence of ferromagnetic iron sulfides (IS). All the remanence is lost at ~580°C corresponding to the Curie temperature of magnetite (Mg). In specimen 2, a decrease in remanence is observed at ~120°C. This could correspond to the Néel temperature of goethite (G) or could be due to the removal of a viscous component.

Figure 4: a) Evolution of the RT-SIRM during a cooling-warming cycle of preserved COx fragments from FOR1118. The increase in remanence during cooling is due to the presence of goethite (G%=25). Note the presence of the Verwey transition of magnetite that is not recovered upon warming. b,c) Evolution of the RT-SIRM on three successive cooling-warming cycles for
powdered FOR1118 COx claystone (b: first cooling-warming cycle, c: second and third cycles) after the heating phase at 395 K under a 2.5 T field. The increase in remanence during cooling between 300 and 10 K is 30, 30 and 25% respectively. d, e) Evolution of the RT-SIRM during a cooling-warming cycle for two unpreserved COx samples. The increase from 300 to 10 K is 46 and 50% for d) FOR1118 sample left for 24h in air after crushing and e) K119 (unpreserved since its collection) respectively.

Figure 5: a) Macroscopic aspect of centimeter-sized pyriteous ammonite from the ANDRA URL gallery; b) Backscattered electron image of framboidal pyrite; c) Example of Raman spectrum acquired on pyriteous ammonite providing evidence of pyrite and excluding iron oxyhydroxide; d) Macroscopic view of organic matter-rich ammonite fossil in the Callovian-Oxfordian clay EST26479 core sample from the FOR1118 borehole – unpreserved sample; e) Backscattered electron image of a pyriteous ammonite from the same sample – unpreserved sample.

Figure 6: Evolution of the RT-SIRM during a cooling-warming cycle for a pyrite+organic matter extract. The irreversibility of the curve upon warming may indicate maghemite occurrence.

Figure 7: Evolution of the LT-SIRM of FOR1118 COx sample upon warming to 300 K. LT-SIRM decreases by 60% from 10 to 50 K, indicating significant amount of small particles.
See supplementary material.

### TABLES

<table>
<thead>
<tr>
<th>Sample name</th>
<th>Type</th>
<th>Location</th>
<th>Formation/ Age</th>
<th>Collection date</th>
<th>Preservation</th>
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<tbody>
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<td>EST26479</td>
<td>Claystones (fragments)</td>
<td>FOR1118 (-124.06 m)</td>
<td>(-)</td>
<td>C2b1 (COx)</td>
<td>2009</td>
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<td>Claystones (powder)</td>
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<td>(-)</td>
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<td>2009</td>
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<tr>
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<td>Claystones (powder)</td>
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<td>(-)</td>
<td>C2b1 (COx)</td>
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<td>(-)</td>
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<td>COx</td>
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<td>Unpreserved</td>
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<td>OM + pyrite (tr.) (powder)</td>
<td>FOR1118 (-124.06 m)</td>
<td>(-)</td>
<td>COx</td>
<td>2009</td>
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Table 1: List of the samples from the Callovian-Oxfordian formation in the Paris Basin analyzed in this study.
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<th>Mineral Phases</th>
<th>Abundance (% mass)</th>
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<tr>
<td></td>
<td>K119&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Carbonates</td>
<td></td>
</tr>
<tr>
<td>Calcite</td>
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</tr>
<tr>
<td>Ankerite</td>
<td>6</td>
</tr>
<tr>
<td>Dolomite</td>
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</tr>
<tr>
<td>Siderite</td>
<td>0</td>
</tr>
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<td><strong>Total</strong></td>
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</tr>
<tr>
<td>Silicates</td>
<td></td>
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<td>Quartz</td>
<td>21</td>
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<tr>
<td>K-feldspar</td>
<td>4</td>
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<tr>
<td>Plagioclase</td>
<td>2</td>
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<tr>
<td><strong>Total</strong></td>
<td><strong>27</strong></td>
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<tr>
<td>Phyllosilicates</td>
<td></td>
</tr>
<tr>
<td>Illite + Illite/smectite</td>
<td>43</td>
</tr>
<tr>
<td>Chlorite</td>
<td>2</td>
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<tr>
<td>Biotite</td>
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<tr>
<td>Kaolinite</td>
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</tr>
<tr>
<td><strong>Total</strong></td>
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<tr>
<td>Others</td>
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<td>Pyrite</td>
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<tr>
<td><strong>Total Organic Carbon (TOC)</strong></td>
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</table>

<sup>a</sup> After Tournassat et al. (2007).

<sup>b</sup> Determined from modeling of XRD patterns and chemical data, following the same procedure than that of K119.

Table 2: Mineralogical phases and total organic carbon present in the two Callovian-Oxfordian claystones studied in this paper.
Table 3: Calculation of pH, pe, Fe concentrations and Fe phases saturation indices (SI) according to the model and the database from Gaucher et al. (2009) and to four different hypotheses on the Fe concentration and redox controls.

<table>
<thead>
<tr>
<th>Systems</th>
<th>pH</th>
<th>pe</th>
<th>Fe (µmol L⁻¹)</th>
<th>SI Goethite</th>
<th>SI Magnetite</th>
<th>SI Pyrite</th>
<th>SI Siderite</th>
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</thead>
<tbody>
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<td>Siderite/pyrite</td>
<td>7.22</td>
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<td>47</td>
<td>1.0</td>
<td>0.9</td>
<td>0.0</td>
<td>0.0</td>
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<td>-2.0</td>
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<td>Magnetite/pyrite</td>
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<td>0.7</td>
<td>0.0</td>
<td>0.0</td>
<td>-0.4</td>
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<tr>
<td>Goethite/siderite</td>
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<td>-3.9</td>
<td>47</td>
<td>0.0</td>
<td>-1.2</td>
<td>13.3</td>
<td>0.0</td>
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<td>-3.9</td>
<td>149</td>
<td>0.0</td>
<td>-1.0</td>
<td>15.0</td>
<td>0.3</td>
</tr>
</tbody>
</table>
Figure 1

Figure 2
Figure 3
Figure 4
Figure 5
Figure 6

Figure 7
Figure 1
Figure 2
Figure 3
Figure 4
Figure 5
Figure 6
Figure 7