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Evolution of porewater composition through time in limestone aquifers: salinity and D/H of fluid inclusion water in authigenic minerals (Jurassic of the eastern Paris Basin, France)

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Abstract

Past water circulations can significantly reduce the porosity and permeability of marine limestones. This is particularly the case in the Middle (Bathonian / Bajocian) to Upper (Oxfordian) Jurassic limestones from the eastern border of the Paris Basin. The knowledge of the timing, the temperature and composition of paleowaters is essential to model the hydrological evolution in this area where the Callovian-Oxfordian claystones are studied for the storage of nuclear wastes. In this way, fluid inclusions hosted in low-temperature (<60 °C) authigenic calcite, quartz and celestite crystals were analyzed by Raman spectroscopy and mass spectrometry to determine the chlorinity and D/H ratios. Chlorinity measurements (mmol Cl per liter of water) in fluid inclusions trapped in authigenic crystals during the late Jurassic / early Cretaceous period revealed unexpected high values, up to 3800 mmol·l⁻¹, indicating that brines were involved in some of the diagenetic crystallization processes. By contrast, fluid inclusions in calcite cements of Cenozoic age within the Oxfordian limestones have low Cl concentration (less than 150 mmol·l⁻¹), thus showing that a dilution event caused by water infiltrations during the Cretaceous uplift of this part of the basin has flushed out the original saline porewater. By coupling δD of fluid inclusion with δ¹⁸O of calcite crystals, we estimate that calcite precipitation occurred at temperatures between 25 and 53 °C. The

hydrogen isotope composition of calcite-forming water is different between the Middle Jurassic (δD ranging from -20 to -35.8 ‰ $V\text{-SMOW}$) and the overlying Oxfordian limestone (δD from -59.5 to -44.8 ‰ $V\text{-SMOW}$). Present-day groundwaters are also of distinct composition on both sides of the Oxfordian claystones, indicating that limestone aquifers underwent independent hydrologic evolutions since the early diagenetic Jurassic cementation.

Keywords: Paris Basin, Jurassic, fluid inclusions, hydrogen stable isotopes, salinity, paleohydrology

Introduction

In the upper part of sedimentary basins, low-temperature diagenetic processes (<60 °C) may significantly modify the original porosity and permeability of the sediments through precipitation of authigenic cements. In such conditions, the chemical and isotopic composition of the fluids involved can be either characterized indirectly (e.g., chemical and isotopic analyses of authigenic minerals) or directly through the study of fluid inclusions (e.g. Shepherd et al., 2000; Sandström and Tullborg, 2009; de Haller et al., 2011). The latter approach is somehow challenging in low-temperature systems. Indeed, fluid inclusions trapped at temperatures lower than ~ 50 °C are generally very scarce, metastable and composed of one phase (liquid) at room temperature (Goldstein, 2001), making impossible accurate microthermometric measurements (excepted by forcing the nucleation of a vapor phase, see Krüger et al., 2011). The evolution of the paleofluid chemistry from ancient hydrologic conditions up to present day is however of critical importance for the understanding of the regime of element mobility in modern aquifers (Hendry et al., 2015).

This is particularly the case for the aquifer-aquiclude system of the eastern part of the Paris basin, for which models of fluid circulation are needed. Indeed, since 1994, the French national radioactive waste management agency (Andra) has been studying the feasibility of a long-term geological disposal of intermediate to high level long-lived nuclear wastes. In this framework, an underground research laboratory (URL) was built in the East of the Paris Basin (Meuse/Haute-Marne) in a 150 m thick succession of Callovian - Oxfordian (COx) clay-rich rocks (present day depth between ~ 400 and 550 m). This formation is sandwiched between two limestone-dominated units: the overlying Oxfordian to Tithonian (Upper Jurassic) and the underlying Bathonian and Bajocian (Middle Jurassic) formations. Exchanges of gas and solutes through the Callovian-Oxfordian claystones occurred by diffusion (Lavastre et al., 2005), this process being responsible of the entire replacement of the original porewater at the Ma scale (Giannesini, 2006). Hence, the characterization of geological paleofluids related to the main fluid events helps understanding present porewater composition.

Both Oxfordian and Middle Jurassic limestones have been intensively cemented by ancient water circulations (Clauer et al., 2007; Brigaud et al., 2010; Lavastre et al., 2011). With the exception of early cementation in phreatic or vadose environments, most of the porosity is filled by successive stages of calcite spars (Buschaert et al., 2004; Vincent et al., 2007; Brigaud et al., 2009a; Carpentier et al., 2014).

Analyses were performed during the past decade on calcite cements, particularly oxygen and carbon stable isotope data (Buschaert et al., 2004; Hibsich et al., 2005; Vincent et al., 2007; Brigaud et al., 2009a; Carpentier et al., 2014), strontium isotope data (Maes, 2002; Hibsich et al., 2005; Brigaud et al., 2009a), and Rare Earth Elements (Carpentier et al., 2014). These studies highlighted several stages of calcite cementation inferred to Early Cretaceous uplift

phases and Cenozoic tectonic stress periods. According to their $\delta^{18}\text{O}$ values and considering temperatures below 60 °C, Early Cretaceous calcite cements have likely originated from mixed marine and meteoric waters, whereas Cenozoic cementations are of meteoric origin (Buschaert et al., 2004; Brigaud et al., 2009a; Carpentier et al., 2014). Cenozoic cementation in the Oxfordian limestones filled most of the pore space and constitutes the only macroscopic crystals in geodes and vugs. Despite numerous available isotopic data on these cements, the precise origin of the calcite-forming water remains uncertain, as the temperature of crystallization derived from only six homogenization temperatures of primary fluid inclusions in calcite cements within the Oxfordian. Buschaert et al. (2004) estimated trapping temperatures between 32 and 42 °C after pressure correction.

Although somehow challenging, the measurements of fluid inclusion stable isotope compositions (δD and $\delta^{18}\text{O}$) might be of primary importance in the discussion of fluid-rock interactions (e.g., Mazurek, 1999; Tarantola et al., 2007; Dublyansky and Spötl, 2010). However, with the exception of the preliminary study of Buschaert et al. (2004), no chemical and isotopic analyses of fluid inclusions were carried out and thus, the paleohydrological evolution of the studied area is still not fully understood (Dublyansky, 2004).

This paper presents a detailed investigation of fluid inclusions trapped in calcite, celestite and quartz cements in the Middle to Upper Jurassic limestones and marls formations in the eastern Paris Basin. It explores 1) how the analysis of fluid inclusions trapped in authigenic crystals provide information regarding the composition and temperatures of paleo-groundwaters in a low-buried sedimentary basin, and 2) how to reconstruct the timing and origin of fluid circulations in relation to regional geodynamic events from deposition time to present.

Geological setting

The intracratonic Meso-Cenozoic Paris Basin was initiated after a general collapse of the Variscan massif during Permian times (e.g., Mégnien et al., 1980). A westward shift of the subsidence towards the present day center of the Basin occurred during late Triassic (Carnian), which led to the observed monocline structure. The eastern Paris Basin sedimentary formations gently dip to the West and show only local evidences of deformation. Triassic sediments consist of fluvial siliciclastic deposits (Buntsandstein facies) grading into shallow-water marine carbonates (Muschelkalk facies) and evaporites (Keuper facies). The Lower Jurassic sedimentary formations consist mainly of marls and shales deposited during the Carnian to Toarcian stratigraphic cycle (Guillocheau et al., 2000). This period corresponds to a large opening of the Basin and to a transgression of the Tethys sea from the East. The early Bajocian marked the transition to a vast carbonate environment (Durlet and Thierry, 2000), with the deposition of oobioclastic sediments together with coral buildups. A major change of facies occurred at the early / late Bajocian transition, with mixed carbonate (oid-dominated) and siliciclastic sedimentation (Brigaud et al., 2009b). A new carbonate ramp then developed in the northeastern Paris Basin during the Bathonian, with typical inner ramp depositional environment (oolitic shoal or lagoon). A general drowning of the platform resulting in deposition of clay-rich sediments started during the Callovian and lasted until the early Oxfordian. The depositional environment gradually returned to carbonate sedimentation during the Middle Oxfordian, with reefal-dominated and oolitic limestones. Lowstand relative sea-level favored the increase of siliciclastic inputs at the beginning of the late Oxfordian (Carpentier et al., 2010). A major carbonate crisis occurred at the Oxfordian/Kimmeridgian transition (Lefort et al., 2011), with marl dominated

sedimentation, coming next with a mixed carbonate and siliciclastic ramp environment during the Tithonian. During late Cimmerian and late Aptian, two stages of emersion associated with erosion occurred (Quesnel, 2003), as a result of both eustatic sea-level drop and distant influence of uplift of rift shoulder linked to the North Atlantic rifting (Ziegler, 1990). A large transgression occurred in late Cretaceous times resulting in the deposition of chalk. The eastern basin margin definitively emerged and experienced weathering and erosion during the Cenozoic, allowing the exhumation of the underlying Jurassic and Triassic sediments, which presently crop out in the area.

The tectonic agenda in this part of the Basin was summarized by André et al. (2010). Two main stages have been distinguished:

i) Early stages, as recorded by the formation of discrete clay minerals (I/S minerals dated around 150 ± 10 Ma (Rousset and Clauer, 2003). Several episodes of thermal anomalies linked to the rifting of the central Atlantic Ocean are well known in Western Europe. Major hydrothermal events occurred during the Jurassic (190-170 Ma and ~ 150 Ma) resulting in Pb-Zn-Ba-F and U mineralizations along the margins of the basin, and in the Armorican, Central and Morvan massifs (Lancelot et al., 1984; Bonhomme et al., 1987; Respaut et al., 1991; Cathelineau et al., 2004, 2012), as well as widespread illitization in Permian and Triassic sandstones (Clauer et al., 1995). Other major extensional events favoring fluid circulations are known at ~ 120 Ma and 80 Ma (Clauer et al., 1995, 1996). All these events coincide with an increase in the subsidence rate in the Paris Basin (Ferry et al., 2007).

ii) Main tectonic events occurred during the Cenozoic, as the basin was submitted to the Pyrenean and Alpine orogenies, as well as extensional movements symmetric and synchronous of the continental rifting of the Rhine graben, such as the Gondrecourt graben delimiting the URL area (Fig. 1). Major faults, inherited from the Hercynian orogeny may also have been active during the Mesozoic and then reactivated by the Pyrenean and Alpine events (André et al., 2010).

The geothermal gradient in the URL area is around $3 \text{ }^\circ\text{C} / 100 \text{ m}$ (Landrein et al., 2013). The burial and thermal history of the Mesozoic series was reconstructed by the study of several independent geothermometers (Blaise et al., 2014). Maximum burial temperatures, reached during the Late Cretaceous, were estimated of $40 \pm 5 \text{ }^\circ\text{C}$ in the Upper Oxfordian and $60 \pm 5 \text{ }^\circ\text{C}$ in the Middle Jurassic limestones (Fig. 2).

Sampling

Core samples

The macroscopic evidences of water-rock interactions during burial diagenesis and telogenesis are the diagenetic/hydrothermal cements that may fill the porous space in limestones located on both sides of the Callovian-Oxfordian claystones. These evidences are scarce, but vugs and geodes were found at the base of the Oxfordian limestones, and within the Bajocian formation. Samples have been recovered thanks to five drill cores cutting through all the series near the URL site (Fig. 1). Seven samples hosting vugs were selected for D/H analysis of water trapped within fluid inclusions (four in the Oxfordian and three in the Middle Jurassic limestones, Fig. 1, Table 1). Samples consist of euhedral calcite, either scalenohedral or globular, found in vugs originating from bioherm units, and of quartz-celestite assemblage found in cavities originating from the dissolution of bioclasts and compaction cracks in the *Terrain à Chailles*, at the top of the Callovian-Oxfordian claystones. According to U-Pb dating, authigenic calcite crystallized at two distinct periods (Pisapia et al.,

2011): 149.2 ± 5.8 Ma in the Middle Jurassic limestones (*i.e.*, during the late Jurassic / early Cretaceous period) and 33.2 ± 5.5 Ma in the Oxfordian (*i.e.*, during the late Eocene – Oligocene period). The authigenic calcite found in the Dogger samples corresponds to the “Bc1” or “Bc2” stages described by Brigaud et al. (2009a), (named “Cal1” and “Cal2” in Carpentier et al., 2014). The samples selected for D/H measurements were also prepared for fluid inclusion petrography and salinity measurements (Table 1). Four other samples were investigated for the measurement of the salinity of water of fluid inclusions by Raman spectroscopy in quartz (one sample in the Lower Oxfordian) and celestite (three samples in the Middle Oxfordian, Lower Oxfordian and Bathonian) (Table 1). Other core samples from Lower Oxfordian to Bathonian series were selected for mineral isotopic analyses (O, C, Sr) (Table 2). As sampling was carried out from five different wells in the Andra studied area, the depth of each sample is normalized to the depth measured in the borehole HTM102 (Fig. 1D).

Outcrop samples

Seven outcrop samples have been selected (Fig. 1, Table 1): five fracture-filling calcite grains in hydrothermal breccias formed during the Oligocene, crosscutting the Upper Oxfordian to Lower Kimmeridgian limestones, at the eastern flank of the Gondrecourt graben (Augeville), one fracture-filling calcite, also formed during the Oligocene, within the Upper Oxfordian limestones near Leurville and one vug-filling calcite in the Bajocian limestones sampled at the Sommerécourt quarry, close to the “Vittel fault”.

Analytical procedures

1. Petrography

Conventional 30 μm thin sections were prepared for each sample for petrographic observations. All samples were observed by cathodoluminescence, with a 8200 CCI CITL apparatus, coupled with an Olympus microscope provided with an AxioCam digital camera (*GeoRessources* laboratory, Nancy, France). In order to provide complementary information on calcite composition, some thin sections were stained after reaction with an alizarin-potassium ferricyanide solution. Fluid inclusion petrography and Raman spectroscopy analyses were carried out from double-polished thick sections (200 – 300 μm).

2. Raman spectroscopy

Fluid inclusions in quartz, celestite, and calcite were investigated with a LabRAM HR Raman spectrometer for gas and chlorinity (chloride concentration) determination, using a monochromatic beam at 457 nm, at the *GeoRessources* laboratory (Nancy, France). In calcite, the efficiency of this method can be limited by several factors, including (1) the size and shape of fluid inclusions (predominantly < 5 μm and flat shaped), (2) high birefringence of calcite and (3) fluorescence of the mineral matrix. The latter parameter strongly depends on the amount of Fe^{2+} in calcite. Indeed, ferroan calcite usually shows some fluorescence, which may prevent any spectroscopic characterization of the fluid phase. Fluid inclusion chlorinity was obtained from the stretching band of water at mineral optical extinction, following the methodology described in Dubessy et al. (2002), Baumgartner and Bakker

(2008), Caumon et al. (2013, 2015) and Tarantola and Caumon (2015). Results are given in $\text{mmol}\cdot\text{l}^{-1}$ with an accuracy of $\pm 10\%$.

3. Isotopic measurements

The hydrogen and oxygen isotopic signatures (δD and $\delta^{18}\text{O}$) of paleowaters trapped within fluid inclusions are obtained by two different ways. The δD value is obtained by direct analysis of the water after extraction from the host-mineral. Because we are dealing with primary fluid inclusions, calcite, celestite and quartz precipitated from the circulating fluid. The $\delta^{18}\text{O}$ value of the paleofluid is obtained from the oxygen isotopic value of the host-mineral after equilibrium fractionation calculation at selected temperatures.

Hydrogen isotopes

Calcite and celestite crystals were weighted (1.0 to 8.2 g, Table 1) and introduced into steel tubes. Calcite-bearing tubes were connected to a vacuum extraction line and heated overnight at $120\text{ }^\circ\text{C}$ to release H_2O molecules adsorbed at the mineral surface (Dublyansky and Spötl, 2009). Preliminary tests conducted under the microthermometric heating stage had shown that our fluid inclusions did not decrepitate, nor stretch, for temperatures below $200\text{ }^\circ\text{C}$. The calcite-bearing tubes were subsequently crushed under vacuum to release all fluids hosted in crystals. H_2O (and possibly CO_2) were trapped under vacuum in a liquid nitrogen-cooled U-shaped tube, incondensable gases being directly removed from the extraction line. CO_2 was never found in sufficient amount to be analyzed, and was thus removed from the line. H_2O was reduced into H_2 by passing through a uranium reactor heated at $800\text{ }^\circ\text{C}$. H_2 was trapped in glass tubes and analyzed using a VG 602D mass spectrometer at CRPG laboratory (Nancy, France). All results are expressed with respect to V-SMOW. Two in-house standards were weekly extracted and analyzed to optimize the accuracy of the isotopic measurements.

An advantage of the metal reduction method for the preparation of gas sample for hydrogen isotope ratio analysis is the small size of the sample $< 1\ \mu\text{l}$ (Wong et al., 1984). Water is reduced into H_2 by passing through a uranium reactor heated at $800\text{ }^\circ\text{C}$ (Bigeleisen et al., 1952). The same reagent is used for the reduction of consecutive water samples. Because water adheres to the internal surfaces of the vacuum system, D/H isotope analysis experiences a memory effect, where the measured D/H ratio of a sample is displaced, from its true value, towards the D/H ratio of the previous samples. Memory effects are most pronounced when consecutive samples have substantially different isotopic compositions as well as when the amount of water reduced in the reactor is significantly different. To avoid this issue, samples were systematically duplicated, and the δD deviance between each isotopic measurement was less than $\pm 1.5\text{ ‰}$. For each duplicate, δD values from the first extraction were thus not considered, as potentially affected by this memory effect. Taking into account the overall reproducibility of internal standards and sample duplicates, the accuracy of D/H measurements is lower than 3 ‰ .

C and O stable isotopes

Calcite spars, quartz and amorphous silica were extracted from fractures and vugs and picked out under the microscope to select monogenic crystals. The finely crushed carbonate and silicate materials were reacted respectively, with anhydrous H_3PO_4 at $50\text{ }^\circ\text{C}$ during 15

hours and with BrF_5 in Ni tubes at 670 °C overnight (following the method of Clayton and Mayeda, 1963). After silicate dissolution, the liberated O_2 was converted into CO_2 by reaction with hot graphite. Isotopic analyses were carried out on CO_2 gas using a VG SIRA 10 mass spectrometer at *Geosciences Rennes* laboratory (France) and expressed with the conventional delta notation vs. V-SMOW (O) and PDB (C). Analytical precision was quantified at ± 0.1 ‰ for both C and O, using in-house carbonate standard *Prolabo Rennes* and NBS 19 reference material.

Sr isotopes

Picked celestite crystals were partially dissolved in HNO_3 . Separation of Sr for isotopic determination was conducted on a Sr-resin, following the methodology described by Pin et al. (2003). Leachates were evaporated, redissolved in HNO_3 and deposited on Ta filaments. Rb and Sr isotopes were measured on a Triton TI TIMS at *GIS laboratory* (Nîmes, France). Values are reported as $^{87}\text{Sr}/^{86}\text{Sr}$ ratios. The external reproducibility of the isotopic measurements was controlled by periodic analysis of the NBS 987 standard providing a mean $^{87}\text{Sr}/^{86}\text{Sr}$ ratio of 0.710258 ± 2.10^{-6} (2σ). The $^{87}\text{Sr}/^{86}\text{Sr}$ ratios were measured at least 90 times to ensure an analytical error below 5.10^{-6} (2σ).

Results

1. Petrographic description

Crystallization of authigenic minerals can be macroscopically observed within vugs and compaction cracks (Fig. 3a, b). Euhedral sparite fringe formed at the border of the micritic matrix ("Spar 1", Fig. 3c, d, e, f). Their coloration by reaction with an alizarin solution revealed a clear gradation in composition, with an increase of the Fe/Mg ratio towards the edge of the crystals (Fig. 3f). They are followed by amorphous silica, which forms spherulites or irregular fringes (Fig. 3c-h), sometimes crosscutting calcite spars (Fig. 3f). Quartz postdates chalcedony and occurs either as microcrystalline fringe (Fig. 3c, d) or euhedral crystals (Fig. 3f, g). The remaining space is filled by celestite (Fig. 3c, d, f) or sparite (Fig. 3e). The second stage of sparite ("Spar 2") crosscuts, and thus postdates, celestite infillings (Fig. 3h). A detailed petrographic description of the calcite samples can be found in Carpentier et al. (2014) and reference therein.

2. O, C and Sr isotopes

O and C stable isotope measurements were carried out for calcite samples, O stable isotopes for quartz and amorphous silica and Sr stable and radiogenic isotopes for celestite crystals. All results are presented in Tables 1 and 2. The O isotope composition ($\delta^{18}\text{O}$) of calcite from the vugs is between 21.4 and 24.2‰ and between 20.7 and 21.9‰ for the hydraulic breccias of the Upper Oxfordian (locality of Augeville and Leurville). All $\delta^{13}\text{C}$ values are positive, between 0.0 and +3.1‰. In the boreholes (Table 2), the calcite spar has comparable $\delta^{18}\text{O}$ and $\delta^{13}\text{C}$ values of 21.2 and +1.5‰, respectively. Quartz and chalcedony have $\delta^{18}\text{O}$ values of 33.2 and 32.4‰, slightly above the calcite value of the host marl ($\delta^{18}\text{O} = 30.0$ ‰).

3. Fluid inclusions

Petrography

As mechanical crushing of crystals releases the bulk fluid content trapped within the inclusions, careful petrography was carried out on fluid inclusions in calcite crystals (Fig. 4). Fluid inclusions are typically of small size ($< 5 \mu\text{m}$) and exhibit a single metastable aqueous liquid phase at room temperature (Fig. 4a, b), which prevents the determination of fluid salinity and minimal trapping temperatures by standard microthermometric methods. No nucleation of a vapor phase occurred within the inclusion after successive cycles of freezing and melting. In calcite, the discrimination between the primary or secondary origin of fluid inclusion cannot always be achieved as both may be aligned along the cleavage planes of the crystals. When fluid inclusion planes are restricted to a single crystal (as illustrated on Fig. 4c), the inclusions can be interpreted either as primary or as pseudo-secondary (Goldstein and Reynolds, 1994). However, this petrographic criteria may be lacking, so the origin of the inclusions remains unclear (Fig. 4b, d). It is worth noting that no clear secondary fluid inclusion planes, crosscutting several crystals, were observed. As a result, the fluid observed in the crystals is assumed to be the fluid present during crystal growth.

Fluid inclusion composition

Raman spectroscopy was performed on the aqueous phase only. Most of the fluid inclusions are free of volatile species with the exception of small amounts of CH_4 and CO_2 rarely detected.

The deformation of the stretching band of water can be assigned to increasing chlorinity values (Dubessy et al., 2002; Baumgartner and Bakker, 2008; Caumon et al., 2013, 2015). Tarantola and Caumon (2015) observed a slight effect of metastability (metastable one-phase (L) versus stable two-phase (LV)) increasing with the degree of metastability of the inclusion. According to the estimated trapping conditions at temperatures lower than $60 \text{ }^\circ\text{C}$ in the H_2O -NaCl system, the negative pressure prevailing within fluid inclusions should lie in the range 70-100 MPa at Raman spectroscopy measurement conditions ($20 \text{ }^\circ\text{C}$). In this case, metastable fluid inclusions record a salinity overestimated of maximum 5 to 8 relative mass% NaCl compared to what would be measured in a stable two-phase fluid inclusion (Tarantola and Caumon, 2015).

Both core and outcrop samples of authigenic calcite within the Oxfordian limestones, as well as samples from the Bajocian quarry at Sommerécourt, are characterized by low to non-saline fluid inclusions (Cl concentration between 0 and $160 \text{ mmol}\cdot\text{l}^{-1}$, Table 1). By contrast, vug-infillings calcite from the Middle Jurassic core samples contains a much more saline fluid. These brines have a chlorine concentration ranging from 650 to $2700 \text{ mmol}\cdot\text{l}^{-1}$. Such values are significantly higher than the average present-day seawater ($\sim 550 \text{ mmol}\cdot\text{l}^{-1}$, Fontes and Matray, 1993a). Fluid inclusions hosted in quartz and celestite crystals in the Middle to Lower Oxfordian limestones display chlorine concentrations ranging from 320 to $3660 \text{ mmol}\cdot\text{l}^{-1}$ (Table 1). In the Bathonian, vug-filling celestite crystals contain fluid inclusions also with a large range of chlorinity values, between 760 and $3840 \text{ mmol}\cdot\text{l}^{-1}$.

δD of fluid inclusions

Core samples

The hydrogen isotope ratio, expressed as δD , of the water trapped in fluid inclusions within authigenic calcite crystals range between -20 and -60 ‰ (Table 1). In the Oxfordian limestones, the δD values of calcite inclusion-water range between -45 and -60 ‰. In the Middle Jurassic limestones, inclusion water shows by contrast higher δD between -20 and -36 ‰. The hydrogen isotope value of the water trapped in fluid inclusions within celestite is -94 ‰ (Table 1).

Outcrop samples

Hydrothermal breccias formed during the Cenozoic have been sampled in Augeville and Leurville (Fig. 1). They contain numerous fluid inclusion and yield δD values ranging from -40 to -54 ‰ (Table 1). Fluid inclusions trapped in vug-filling calcite in Middle Jurassic limestones from Sommerécourt have comparable δD values: two values were obtained, yielding δD of -48 and -51 ‰.

Discussion

1. Chalcedony-quartz-celestite assemblage as a record of the evolution of porewater composition

The chalcedony-quartz-celestite succession, infilling moldic porosity and compaction cracks are essentially found in the *Terrain à Chailles* silty-marl layer. This formation, usually characterized by its abundant chert nodules, is situated at a lithologic transition between the Callovian-Oxfordian claystones (COx) and the Oxfordian limestone aquifer. In minor amounts, silica and celestite are also found as diagenetic phases within the COx itself (Lerouge et al., 2011), and a single occurrence of celestite has been detected in the underlying Bathonian limestones (this study). The chalcedony-quartz-celestite succession is sometimes predated by pyrite, dolomite and siderite, interpreted as early diagenetic on the basis of petrographic criteria and geochemical analyses (Lerouge et al., 2011). However, no explanation has been so far advanced for the origin of siliceous cements and their specific occurrence at the top of the COx formation. The *Terrain à Chailles* is overlaid by another marly formation, the *Marnes des Eparges*, with a decametric spongolite facies. In this layer, abundant sponge spicules are recrystallized in calcite. The dissolution of biogenic silica is common during early diagenesis (Hendry and Trewin, 1995; DeMaster, 2004; Madsen et al., 2010), SiO_2 being transported in solution and further crystallized as amorphous silica (chalcedony) and quartz. The transition from the amorphous polymorph to euhedral quartz can be gradual (Fig. 3a, b), and is rather due to relative rates of nucleation and growth than the result of a temperature increase (Williams et al., 1985). Using the fractionation coefficient of Kita et al. (1985) between amorphous silica and H_2O , the calculated $\delta^{18}\text{O}$ of parent-water ranges between -1.5 and $+0.9$ ‰, at 30 and 40 °C, respectively. Within the same temperature conditions, $\delta^{18}\text{O}$ of water involved in quartz crystallization ranges from -1.5 and $+1.1$ ‰ (Zheng, 1993). These ranges match the seawater domain (Fig. 5). The very D-depleted water trapped within fluid inclusion of celestite samples possibly records the involvement of biogenic methane during early diagenetic processes. Indeed, methane, either of biogenic or thermogenic origin, is strongly depleted in D (Schoell, 1980). Fluid inclusions with δD ranging from -40 to -110 ‰ have already been documented in various minerals

(calcite, quartz, fluorine, and galena) crystallized at temperatures of 70 to 150 °C, and interpreted as a possible interaction with organic fluids (Charef and Sheppard, 1987; Munoz et al., 1999; Tarantola et al., 2007).

The precipitation of sulfate minerals (gypsum or anhydrite) prior the chalcedony stage is suggested by the preservation of lamellar crystal pseudomorphs (Fig. 3c). Dissolved gypsum or anhydrite associated with chalcedony has been reported in several studies, either in geodes (Boyce et al., 1990; Chowns and Elkins, 1974; Gomez-Alday et al., 2002), evaporitic beds and nodules (Milliken, 1979; Ulmer-Scholle et al., 1993) or cherts (Chowns and Elkins, 1974; Folk and Pittman, 1971; Geeslin and Chafetz, 1982). The precipitation of gypsum or anhydrite likely occurred from sulfate-rich dense brines (Hanor, 2004) percolating downwards, from the surface down to the permeability transition between claystones and limestones. Regarding the large amount of Sr needed to crystallize celestite, the most likely reservoir is coral aragonite (Finch and Allison, 2003), Sr being released during the recrystallization of aragonite into calcite (Baker and Bloomer, 1988; Hoareau et al., 2010). Corals build-up consistently forms several decametric levels within the Middle and Upper Oxfordian limestone units.

The contribution of brines is supported by chlorine concentration of fluid inclusion in quartz and celestite, ranging from 320 to 3620 mmol·l⁻¹, and from 850 to 3690 mmol·l⁻¹, respectively (Table 1). This brine was likely generated by the evaporation of seawater beyond gypsum saturation. The large range of chlorinity provides evidence of a mixing trend between this evaporitic end-member, which has been more or less diluted by seawater and/or meteoric water. Additionally, it is known that Sr concentration in water increases significantly with salinity (Hanor, 2000; 2004). A contribution of secondary brines from the dissolution of Upper Triassic halite is unlikely, regarding ⁸⁷Sr/⁸⁶Sr values of celestite close to Upper Jurassic seawater (Jones et al., 1994), excepting a single value (sample PPA- 360 m, ⁸⁷Sr/⁸⁶Sr = 0.707545, Table 2). A contribution of ascending Triassic fluids would increase the ⁸⁷Sr/⁸⁶Sr ratio, from the dissolution of halite and the interaction with siliciclastic series. Late calcite, postdating quartz (Fig. 3g) and celestite (Fig. 3h), has a δ¹⁸O value of 21.2 ‰. Considering a range of crystallization temperature between 30 and 40 °C (Blaise et al., 2014), δ¹⁸O of parent-water is between -8.8 and -6.5 ‰ (Zheng, 1999), indicating a meteoric origin. Therefore, the chalcedony-quartz-celestite-calcite assemblage provides a unique record of the transition in the composition of porewater, from the concentration of seawater by evaporation to the meteoric water infiltration (Fig. 5).

According to the geodynamical evolution of the basin, meteoric water recharge could have started since early Cretaceous. Thermal anomalies linked to the opening of the central Atlantic Ocean engendered the general emersion of the north-eastern flank of the Paris Basin at the end of Jurassic (during the “Late Cimmerian Unconformity”, LCU). The sea regression is marked by a sabkha environment, with dolomite and anhydrite precipitation during the late Tithonian (“Purbeckian” facies, Guillocheau et al., 2000). At that time evaporated seawater could have migrated downwards due to its higher density, towards the permeability transition between claystones and limestones; alternatively, lateral migration from the north-eastern border of the basin could be invoked. These brines were involved in the precipitation of silica (chalcedony and quartz) and sulfates (celestite and possibly gypsum/anhydrite), before being progressively flushed out by the percolation of fresh meteoric water during the Cretaceous period. Mixing between seawater and meteoric water could also have induced calcite precipitation in the Middle Jurassic aquifer.

2. Comparison with present-day groundwaters

Hydrogeochemistry of present-day underground waters

The chemical and isotopic signatures of present-day groundwater percolating through the porous levels of Oxfordian and Middle Jurassic limestones have been extensively monitored in and around the Andra studied area (Buschaert et al., 2007; Linard et al., 2011; Rebeix et al., 2011). The stable isotope (δD and $\delta^{18}O$) composition of water reveals a strictly meteoric signature (Giannessini, 2006; Linard et al., 2011; Rebeix et al., 2011). Water residence time was estimated on the basis of geochemical tracers to be of about 10 ka for the Oxfordian aquifer and several hundred thousand years in the Middle Jurassic. This implies a recharge of the aquifers under different paleoclimatic conditions. The extremely low-permeable Callovian-Oxfordian claystones prevent any connection between the aquifers that, as a result, should be considered as isolated hydrological systems (Lavastre et al., 2010). However, chemical exchanges of gas and solutes through the Callovian-Oxfordian claystones occur by diffusion (Lavastre et al., 2005; Battani et al., 2011; Bensenouci et al., 2013; Fourré et al., 2011; Mazurek et al., 2011; Rebeix et al., 2011), this process being responsible of the entire replacement of the original Callovian-Oxfordian porewater (Giannesini, 2006).

The thick Lower Jurassic clay-rich sediments, together with the Keuper halite, constitute an efficient isolation between the Middle Jurassic limestones and the deep siliciclastic Lower Triassic aquifer (Marty et al., 2003). Chlorine concentration and $\delta^{37}Cl$ evolution with depth can be interpreted as the result of diffusive transfers from the Keuper halite through the entire Jurassic sedimentary sequence. Such a process may explain chlorine concentrations in the present-day groundwaters (Rebeix et al., 2011).

Towards the central part of the basin, the deep Middle Jurassic and Triassic groundwaters have significant different chemical compositions. Their salinity gradually increases from fresh or brackish waters to brines in the deepest part of the basin (Fontes and Matray, 1993b; Matray et al., 1994; Pinti et al., 1997; Worden et al., 1999; Millot et al., 2011), as a result of a complex mixing process between both primary and secondary brines together with dilution related to meteoric water inputs (Matray and Fontes, 1990). In addition, cross-formational flow from the Triassic sandstones to the Middle Jurassic limestones may have occurred through the fracture network (Worden and Matray, 1995).

Stable isotope signature of paleowaters

As discussed above, petrographic criteria were not always clear enough to determine accurately the relative age of fluid inclusions trapped in calcite crystals. However, a large majority of fluid inclusions is likely of primary origin, and no clear secondary fluid inclusion planes were identified. Therefore, bulk fluid δD values can be considered as representative of the water involved in calcite precipitation. Previous published studies on δD measurements from fluid inclusions hosted in low-temperature calcite have shown that they preserve the original isotopic composition of the forming waters (McGarry et al., 2004; Zhang et al., 2008; Dublyansky and Spötl, 2009, 2010)

The δD value of water trapped in fluid inclusions is reported as a function of the $\delta^{18}O$ values of calcite parent-fluids in Figure 6. The oxygen isotope equilibrium fractionation between calcite and water ($\Delta^{18}O_{Ca-H_2O} = 28.8\text{‰}$) was calculated considering a mean crystallization temperature of 35 °C, and using the calcite-water fractionation factor of Zheng (1999). The value of 35 °C is used here to normalize all samples and be able to visualize them in a δD vs.

$\delta^{18}\text{O}$ plot. This temperature is close to those calculated from fluid inclusion homogenisation temperatures in authigenic calcite in the Oxfordian limestones (Buschaert et al., 2004), but also to the first and second generation of calcite in the Middle Jurassic limestones (Brigaud et al., 2009a).

With the exception of the Middle Jurassic sample 481 m, the isotopic composition of calcite-forming water plots close to the Global Meteoric Water Line (GMWL, Craig, 1961). This illustrates the predominance of meteoric water within the aquifers. As illustrated on Fig. 6, a crystallization temperature of 35 °C leads to a shift of the isotopic composition of some Oxfordian paleowaters (samples 569 m, 660 m, 134 m and 340 m) to the right of the GMWL. This is likely due to an overestimation of the temperature, which should thus be below 35 °C (Fig. 6). Several other processes may also account for the modification of the isotopic composition of meteoric water, among which (1) subsurface evaporation (e.g., Gonfiantini, 1986; Gibson et al., 1993), (2) ^{18}O -enrichment by fluid-rock interactions (e.g., Sheppard, 1986; Matray and Fontes, 1990; Kharaka and Thordsen, 1992) or (3) mixing with other fluids, including seawater (e.g., Lynn Ingram et al., 1996):

(1) Subsurface evaporation leads to the enrichment of the residual water in both ^{18}O and D. However, the mass of evaporated meteoric water isotopically enriched by evaporation is usually limited and tends to be highly diluted by fresh water as it reaches the saturated zone (IAEA, 2000a; 2000b). Present-day groundwaters flowing through the Jurassic aquifers in the eastern or central part of the Paris Basin are for example not significantly affected by evaporation (Matray et al., 1994; Rebeix et al., 2011).

(2) Isotopic exchange reactions between groundwater and carbonates are well documented (Kharaka and Carothers, 1986). In both studied limestone units, the $\delta^{13}\text{C}$ values of calcite are similar to those of host limestones, suggesting that carbon in the newly-formed calcite was supplied by the dissolution-precipitation processes with the host rock (Buschaert et al., 2004; André et al., 2010). Buffering of pore water by carbonate dissolution may also increase its original ^{18}O content (e.g., Clayton et al., 1966).

(3) Significant interaction of calcite-forming waters with liquid or gaseous hydrocarbons can be discarded. No hydrocarbon accumulations have been evidenced within the Jurassic sediments in the eastern part of the Basin because the too low temperatures reached by hydrocarbon source rocks prevented any oil formation (Blaise et al., 2011; 2014). Furthermore, fluid inclusions in calcite are all one-phase aqueous and almost free of volatile species, with only minor amounts of CO_2 and CH_4 rarely detected.

During the whole Jurassic and part of the Cretaceous period, the eastern Paris Basin was immersed (Mégnyen et al., 1980). Hence, the presence of seawater within the porosity of the Jurassic limestones during this period is plausible. This hypothesis was invoked by Brigaud et al. (2009a) to account for the $\delta^{18}\text{O}$ values of the mesogenetic calcite cements in the Middle Jurassic formations. However, mixing of meteoric water with seawater can only generate a small shift relative to the meteoric water line (e.g., Lynn Ingram et al., 1996), unless one or both compounds is/are strongly enriched during evaporation (Knauth and Beeunas, 1986).

In summary, paleowaters showing a shift to the right of the GMWL can be explained either by an overestimation of the crystallization temperature, which should thus be slightly less than 35 °C, or by an enrichment in ^{18}O through the dissolution of host limestones. The latter process is evidenced at present in the center part of the Paris Basin, where long-term equilibration of groundwater with the Middle Jurassic limestones has led to a mean excess in ^{18}O of about 1.5 ‰ relative to the GMWL (Matray et al., 1994) at temperatures between 55 and 80 °C. The δD values of water trapped in fluid inclusions reveal a clear distinction

between the two studied aquifers (Fig. 6). Oxfordian calcite-forming waters are characterized by δD values ranging from -44.2 to -59.5 ‰, consistent with or slightly higher than the present-day groundwater (Fig. 6). As these waters are of meteoric origin, δD values higher than present-day (-49 to -44 ‰) can be explained by different climatic conditions, and hence different δD values of meteoric waters, at the period of paleowater recharge.

δD values of calcite-forming waters in the Middle Jurassic aquifer are clearly distinct from those obtained from the overlying Oxfordian. The δD values of water trapped in calcite cements of Middle Jurassic samples 569 m and 660 m are close, between -32 and -36 ‰. Such values are consistent with the highest δD values measured in present-day groundwater. Moreover, these waters also display high chlorine concentrations ($650 - 2700$ mmol·l⁻¹) so as they are likely enriched in heavier isotopes by the contribution of evaporated seawater.

Paleowater from Middle Jurassic sample 481 m appears highly enriched in deuterium ($\delta D = -20$ ‰). For the water to lie on the GMWL, $\delta^{18}O$ of water has to be calculated at a temperature of 53 °C, which should thus be considered as the minimal temperature for calcite crystallization. This sample also contains saline water trapped as fluid inclusions (Cl concentration between 1000 and 1700 mmol·l⁻¹), and may thus correspond to mixing between evaporated seawater and meteoric water.

Evolution of pore water salinity

The chlorinity of paleowater trapped as fluid inclusions in authigenic calcite from the Oxfordian is comparable to present-day groundwaters (Fig. 7). Within the Oxfordian limestones, widespread calcite cementation was induced by tectonically-driven circulation of meteoric waters during Cenozoic times (André et al., 2010). At this period, the eastern part of the Paris Basin was already uplifted and the aquifers subjected to meteoric water recharge. Field observations at the vicinity of the Gondrecourt graben (Augeville and Leurville sites) show that the Oligocene E-W extensional regime favored the precipitation of large amounts of calcite under a highly dynamic fluid flow inferred from the presence of hydrothermal breccias. The salinity of these calcite-forming waters is very low (< 150 mmol·l⁻¹), thus showing that brines were absent or highly diluted by fresh waters.

In the Middle Jurassic limestones, by contrast, paleowaters trapped in newly-formed calcite reveal high chlorine content, between 650 and 2700 mmol·l⁻¹. Such high salinity is also found in quartz and celestite-hosted fluid inclusions from the Middle and Lower Oxfordian marls and the Bathonian limestones (Table 1). These ranges of values are higher than the average seawater (~ 550 mmol·l⁻¹) and thus imply a brine component in the calcite-forming waters. This brine is likely of primary origin (*i.e.*, generated by evaporation of seawater, Hanor, 1994) as no cross-formational flow from the Upper Triassic halite to the Middle or Upper Jurassic aquifer has been evidenced in the eastern Paris Basin (Marty et al., 2003; Battani et al., 2011; Rebeix et al., 2011). Dating of geodic calcite by the U-Pb method has revealed a major precipitation event at 149.2 ± 5.8 Ma, *i.e.*, during the Kimmeridgian/Berriasian period (Pisapia et al., 2011). During the late Jurassic times, the eastern Paris Basin was still immersed and subjected to marine sedimentation. The opening of the central Atlantic Ocean led to the general emersion of the eastern flank of the basin at the end of Jurassic (during the “Late Cimmerian Unconformity”, LCU). The sea regression is marked by a sabkha environment during late Tithonian, with the deposition of evaporites (“Purbeckian” facies, Guillocheau et

al., 2000). In a similar way, an increase of the Jurassic limestones porewater salinity may have occurred during this period.

The present-day chlorine concentration of Middle Jurassic groundwaters is ten to a hundred times lower than those measured in fluid inclusions trapped in calcite cements (Fig. 7). This illustrates the evolution of porewater composition: the brine component has been progressively flushed out and diluted by meteoric water. This gradual change could have started after the definitive retreat of the sea at the end of Cretaceous times.

3. Fluid circulation on a regional scale

δD and $\delta^{18}O$ of calcite-forming water from different outcrops are shown on Fig. 8. The paleowater isotopic domains in the Oxfordian and Middle Jurassic limestone aquifers from the Andra site are also plotted for comparison on the same figure. Samples from Augeville and Leurville (Fig. 8) correspond to hydraulic breccias crosscutting the Upper Oxfordian limestones and formed during the E-W Oligocene extensional stage (André et al., 2010). As illustrated on Fig. 8, the calcite parent-water plots close to the GWML at temperatures around 35 °C. Furthermore, the isotopic composition of this water is comparable to the one of calcite-forming water within the Oxfordian at depth, in the Andra site. Hence, these results strongly suggest a common origin for these calcite infillings, which is also supported by the Eocene-Oligocene age obtained on geodic calcite from Oxfordian cores (U-Pb dating at 33.2 ± 5.5 Ma, Pisapia et al., 2011). It should be noted that the mean temperature range of paleowater from Augeville and Leurville (inferred from the oxygen and hydrogen isotopic composition) is between 31 and 41 °C, while the one from Oxfordian paleowater at depth in the Andra site is slightly lower, between 25 and 36 °C, if one considers a strictly meteoric origin with no ^{18}O enrichment from the dissolution of host rock. Water circulation may have been initiated during the opening of the Gondrecourt Graben along the major faults, with a subsequent lateral percolation through the Oxfordian aquifer and associate cooling. Noteworthy is the fact that the isotopic composition of geodic calcite sampled within the Bajocian limestones from the Sommerécourt quarry is clearly distinct from those of the Middle Jurassic at depth in the Andra site. Instead, the calcite-forming water inferred from Sommerécourt outcrops rather matches the isotopic composition of meteoric water which has cemented the Oxfordian limestones (Fig. 8). The Sommerécourt quarry is located at the vicinity of the Hercynian "Vittel fault", which was reactivated during the Oligocene extension (André, 2003). Hence, calcite precipitation in these Middle Jurassic limestones likely occurred during the same period as in the Oxfordian limestones (i.e., during Oligocene). Thus, the isotopic composition of the fluid rather marks the process and the period at which large scale water circulation and correlative calcite precipitation occurred, independently of the age of the host rock.

4. Towards a conceptual model of paleowater circulation through time

Two major crystallization events have been characterized in this study:

- During the Late Jurassic/Early Cretaceous period, evidenced by calcite and celestite crystals in the Jurassic limestones, and by chalcedony-quartz-celestite assemblages in the Middle Oxfordian silty-marl layer (*Terrain à Chailles*).
- During the Late Eocene/Oligocene period, evidenced by calcite crystallization in the Upper Oxfordian limestone.

Although not representing the entire paragenetic sequence, these crystallization events are the most significant in terms of cement volumes (Brigaud et al., 2009a; Carpentier et al., 2014) and correspond to two main fluid events. Figure 9 compares the composition of paleowaters with present-day groundwater in a $[\text{Cl}^-]$ vs. δD diagram. This underlines that, in the Middle Jurassic aquifer, both dilution and cooling occurred from ~150 Ma to present. As the water residence time was estimated to be in the order of several hundred thousand years, it is more likely that chlorine in present-day groundwater is supplied by the diffusion from underlying Triassic and Lower Jurassic units, rather than inherited from the ~150 Ma brines identified in this study. In the Upper Oxfordian limestone aquifer, Figure 9 illustrates that paleowaters from the Late Eocene/Oligocene period were warmer compared to present-day groundwater. This may be due to advective water flow under high fluid/rock ratios conditions during the Cenozoic rifting events. As a summary, two sketches are proposed in Figures 10 and 11 to illustrate the main crystallization events, the circulation of paleowater related to the successive geodynamic events and the evolution of fluid sources through time.

Conclusions

In the eastern part of the Paris Basin, the Callovian-Oxfordian claystones are studied for the long-term storage of nuclear wastes (Underground Research Laboratory, ANDRA). Diffusion of gas and solutes from the underlying sedimentary series has erased the original pore water composition. Knowing the evolution of groundwater composition and temperature in the underlying and overlying limestones, from the past geologic times to the present, is a critical step to model the evolution of water advection in limestone aquifers and diffusion through the claystone units. In this study, we have shown that:

- Brines with chlorine concentrations up to $3800 \text{ mmol}\cdot\text{l}^{-1}$ were involved in calcite cementation within the Middle Jurassic limestones and in calcite-quartz-celestite infillings within the Lower Oxfordian marls. These crystallization events likely occurred during the late Jurassic or early Cretaceous period. By contrast, authigenic calcite crystals in the Oxfordian limestones, crystallized during the Cenozoic times, contain fluid inclusions of lower salinity ($< 150 \text{ mmol}\cdot\text{l}^{-1}$), showing that the brines were flushed out by meteoric water infiltrations during the Cretaceous uplift of the eastern border of the basin.
- The oxygen and hydrogen isotope compositions of paleowater are significantly different between the Oxfordian and the Middle Jurassic limestone aquifers. Regarding the Oxfordian, paleowater dated from the Cenozoic times does not exhibit large difference compared to present-day groundwater. Regarding the Middle Jurassic aquifer, water involved in calcite cementation during the Late Jurassic/Early Cretaceous period was probably warmer ($30 - 55 \text{ }^\circ\text{C}$) and of mixed meteoric and evaporated marine origin.
- At the regional scale, fluid inclusions from several calcite infillings of Cenozoic age show have δD values close to those determined in the Oxfordian aquifer at depth, reinforcing the hypothesis of a common origin. Water was warmer at the vicinity of major faults and may have cooled down during the lateral percolation through the aquifer.

From a methodological point of view, this study highlights the fact that chemical and isotopic analyses of fluid inclusions in low-temperature authigenic minerals, although challenging, are critical to reconstruct the evolution of groundwater composition and temperature through times.

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Figure caption

Figure 1. Simplified geological map of the Paris Basin with detailed views of the study area and location of the studied boreholes and outcrops. A. The Paris Basin and its border Hercynian massifs. B. Geological map of the eastern edge of the Paris Basin with major structural features and location of the studied outcrops (star symbols). C. Andra Underground Research Laboratory (URL), with the location of all studied boreholes (EST, PPA, PAX and HTM). D. Simplified lithostratigraphic column of the Jurassic sedimentary sequence, with the position of all studied samples. White squares symbolize quartz-celestite assemblages, grey circles calcite samples.

Figure 2. Burial history diagram with color-scale temperatures (modified from Blaise et al., 2014). Red lines delimit the sequence of interest for the present study.

Figure 3. Petrographic observations.

A. Hand specimen of a compaction crack filled by quartz and celestite (PAX – 324 m, Middle Oxfordian). B. Vug filled by calcite spars (HTM102 – 141 m). C. Paragenetic sequence: Calcite spar fringes (“Spar 1”), chalcedony, micro-quartz and celestite (sample PAX – 324 m, Middle Oxfordian). D. Same as A., SEM – backscattered electron image. E. Cathodoluminescence observation revealing pseudomorphs of lamellar crystals, likely gypsum or anhydrite (EST205 borehole sample, Lower Oxfordian). F. Paragenetic sequence showing chalcedony crosscutting calcite fringes and euhedral quartz crystals. Calcite is stained with an alizarin solution, and the color gradation from purple (core) to blue (edge) shows the increase of the Fe/Mg ratio during crystallization (EST205 borehole sample, Lower Oxfordian). G. SEM – secondary electron picture showing euhedral calcite spar on euhedral quartz (sample PAX – 380 m, Middle Oxfordian). H. Cross-polarized light microphotograph showing a fissure filled with calcite (“spar 2”) crosscutting both celestite and chalcedony (sample EST413 – 376 m, Middle Oxfordian).

Figure 4. A. Fluid inclusion planes distributed along concentric growth zone, witnessing their primary origin (calcite sample AB7). B. Typical small-sized primary aqueous liquid inclusions in calcite exhibiting a single phase at room temperature (sample PPA – 134 m, Upper Oxfordian). C. Fluid inclusion planes (FIP) parallel or aligned at 120° relative to the cleavage planes of calcite (sample PPA – 134 m, Upper Oxfordian). Note that the fluid inclusion planes do not crosscut the crystal border, indicating a primary origin (circle). D. One-phase aqueous inclusions in a celestite crystal (sample PAX – 324 m, Middle Oxfordian).

Figure 5. $\delta^{18}\text{O}$ values of parent-waters for the paragenetic sequence of sample PAX – 324 m (Middle Oxfordian), calculated from $\delta^{18}\text{O}$ of each mineral, using the equations of fractionation of Zheng (1999) for calcite in bulk claystone and late calcite spar, Kita et al. (1985) for chalcedony, and Zheng (1993) for quartz. Calculations were made assuming crystallization temperatures of 35 ± 5 °C (error bars).

Figure 6. Plot of δD vs. $\delta^{18}\text{O}$ (‰_{V-SMOW}) of calcite-parent waters from Oxfordian and Middle Jurassic core samples. $\delta^{18}\text{O}$ values of water are calculated from $\delta^{18}\text{O}$ of calcite, considering crystallization temperatures at 35 °C (Zheng, 1999). The temperatures of crystallization necessary for the calculated water to lie on the GMWL are indicated. Light blue and deep

blue ellipses symbolized the isotopic domain of present-day groundwater in the Oxfordian and Middle Jurassic aquifers, respectively (Rebeix et al., 2011). GMWL: Global Meteoric Water Line (Craig, 1961); SMOW: Standard Mean Ocean Water.

Figure 7. Ranges of chlorinity values ($\text{mmol}\cdot\text{l}^{-1}$) of present-day groundwater and paleowater trapped as fluid inclusions in the aquifers of the Paris Basin. Seawater (SW) and halite saturation values after Fontes and Matray (1993a). When indicated, the error bars illustrate minimal and maximal values measured on fluid inclusions by a given study, where the thick part represents the statistically more representative values. Numbers in parentheses indicate references as follows: (1) Rebeix et al. 2011; (2) Matray et al., 1994; (3) Worden and Matray, 1995; (4) Millot et al., 2011; (5) Fontes and Matray, 1993b; (6) Demars and Pagel, 1994. FI: Fluid inclusions; PB: Paris basin. Light green color is used for the data of this study; deep green color shows present-day groundwater in the study area; orange color represents data from the central part of the Paris Basin.

Figure 8. Plot of δD vs. $\delta^{18}\text{O}$ (‰ $v\text{-SMOW}$) of calcite-parent water from vug and fracture samples within outcropping Oxfordian and Middle Jurassic limestones. $\delta^{18}\text{O}$ values of water are calculated from $\delta^{18}\text{O}$ of calcite, considering crystallization temperatures at 35 °C (Zheng, 1999). The temperatures of crystallization to plot the water along the GMWL are indicated. Light blue and deep blue ellipses symbolized the isotopic domain of present-day groundwater in the Oxfordian and Middle Jurassic aquifers, respectively (Rebeix et al., 2011). Light blue and deep blue parallelograms symbolize the isotopic domain of paleowater determined from core samples of Oxfordian and Middle Jurassic limestones, respectively. A maximum deviation of 1.5 ‰ from the equilibration of water with host limestones is assumed. GMWL: Global Meteoric Water Line (Craig, 1961).

Figure 9. Plot of $[\text{Cl}^-]$ ($\text{mmol}\cdot\text{l}^{-1}$) vs. δD (‰ $v\text{-SMOW}$) measured in fluid inclusions trapped in authigenic calcite and celestite, defining the domain of composition of paleowaters. Arrows show the evolution of porewater composition from the crystallization period to the present.

Figure 10. Conceptual model showing the origin of brines identified in fluid inclusions trapped in authigenic quartz, celestite and calcite crystals in the Oxfordian and Middle Jurassic limestones. The upper part of the figure shows a cross-section of the Paris Basin (not to scale), and major geodynamical events during the Kimmeridgian-Berriasian period.

Figure 11. Conceptual model showing the origin of waters identified in fluid inclusions trapped in authigenic calcite crystals during the Oligocene. The upper part of the figure shows a cross-section of the Paris Basin (not to scale), and major geodynamical events. At that time, the sedimentary series were already dipping

Table caption

Table 1. Oxygen and carbon isotope composition of minerals and fluid inclusion isotopic data and chlorinity in authigenic crystals from core (upper part of the table) and outcrop samples (lower part). For some samples, the $\delta^{18}\text{O}$ value of calcite has been measured on one fraction and the obtained value was applied to the other fractions (in which case “id.” is indicated). The D/H ratio of sample labeled “481 m” was measured twice: in this specific case, the second sample “481 m -2” is a duplicated measurement (“dupl.”). Symbol “/” indicates all other missing values (analyses not performed). $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ values correspond to $\delta^{18}\text{O}$ of parent-waters considering crystallization temperatures of 35 °C using the fractionation equation for calcite-H₂O of Zheng (1999). The range of chlorinity values is indicated with the number of analyses within parentheses.

Table 2. Oxygen, carbon and strontium isotope compositions of authigenic minerals measured on core samples. $\delta^{18}\text{O}_{\text{H}_2\text{O}}$ values correspond to $\delta^{18}\text{O}$ of parent-waters considering crystallization temperatures of 35 °C using the fractionation equation for calcite-H₂O of Zheng (1999). These samples were in too small quantities for the chlorinity and isotopic study of fluid inclusions.

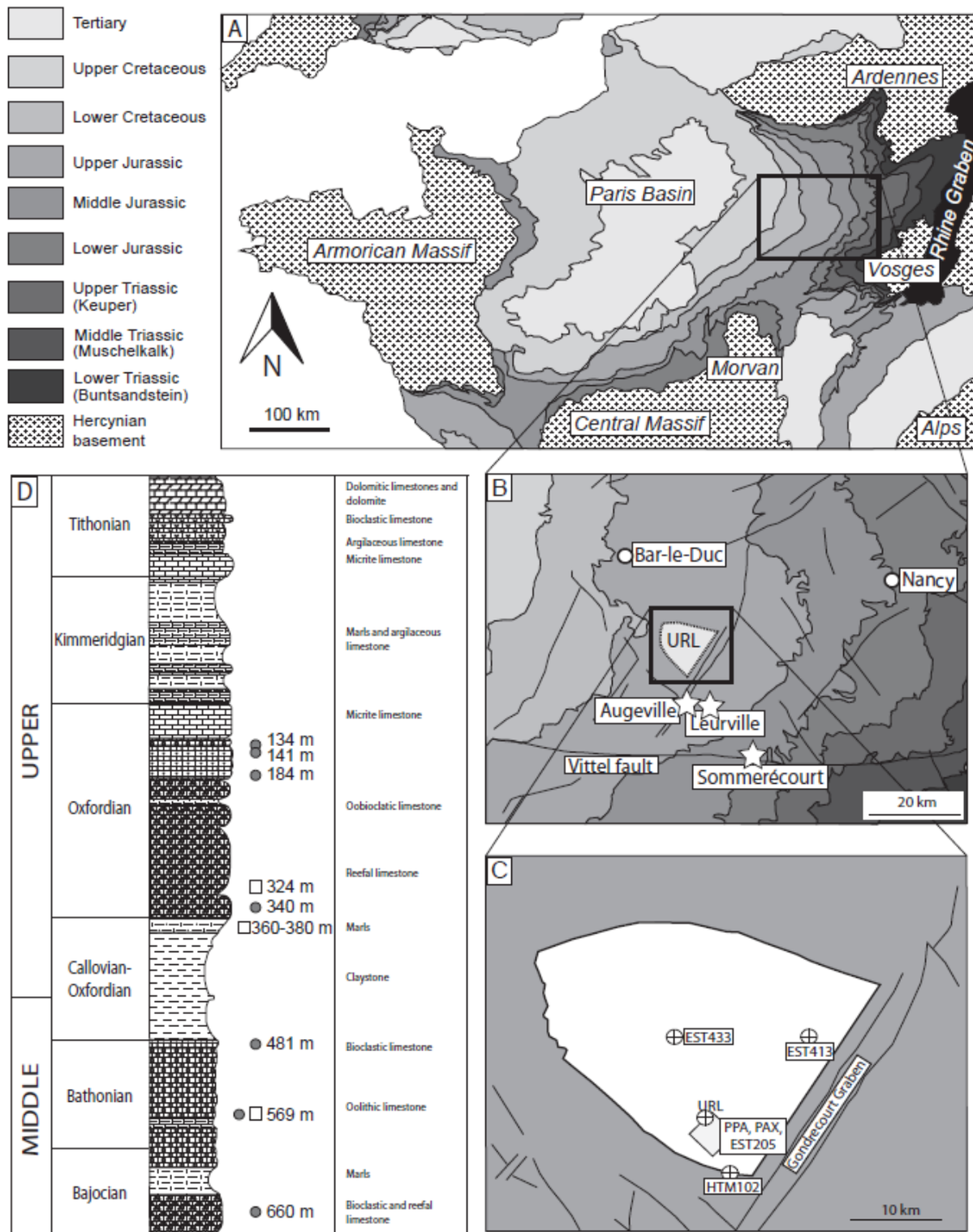


Fig. 1

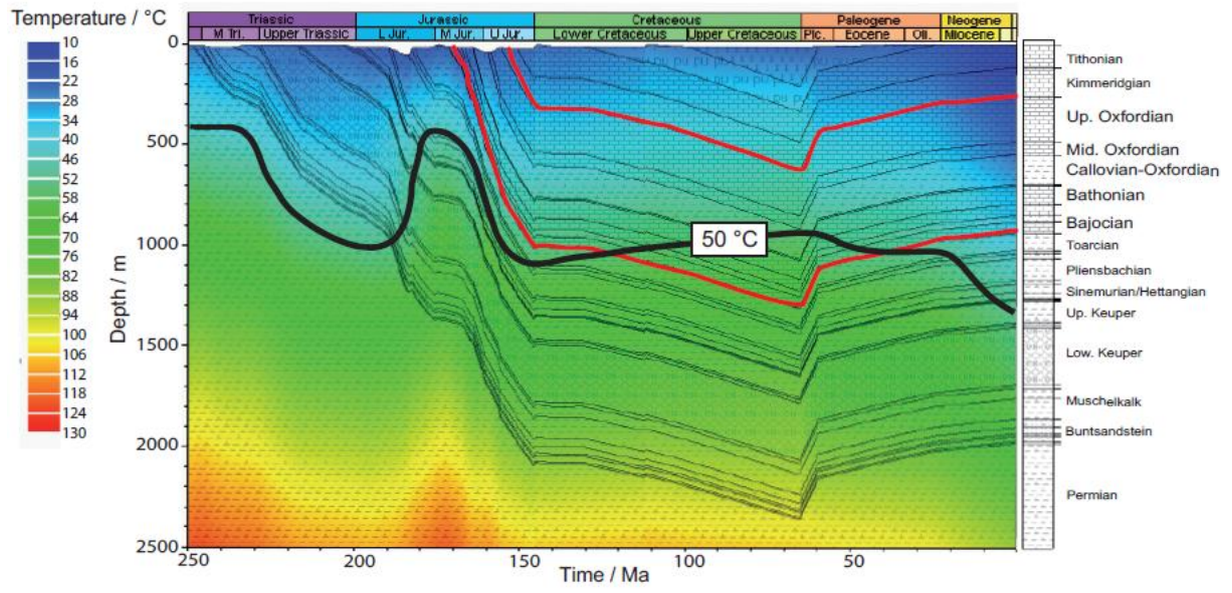


Fig. 2

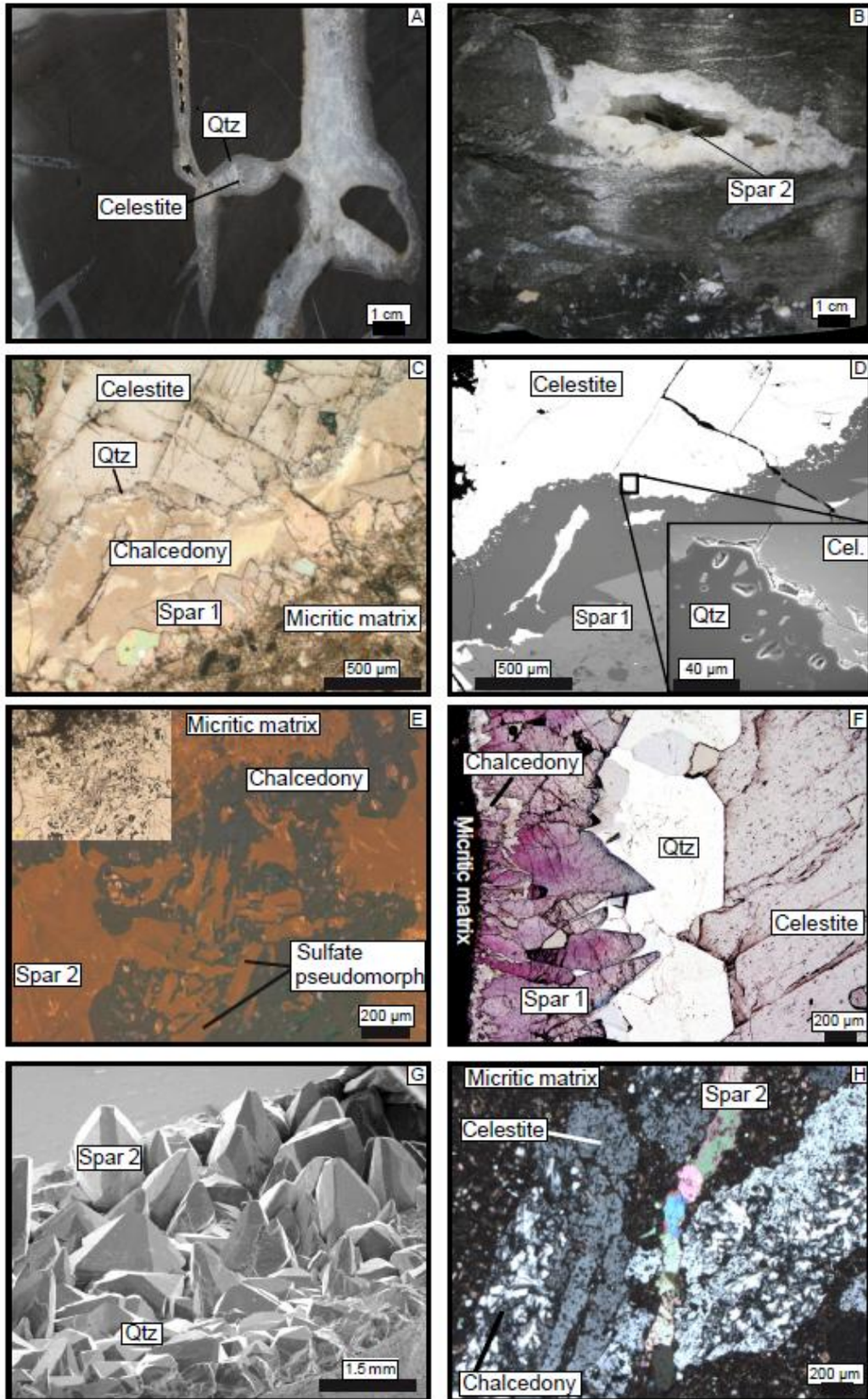


Fig. 3

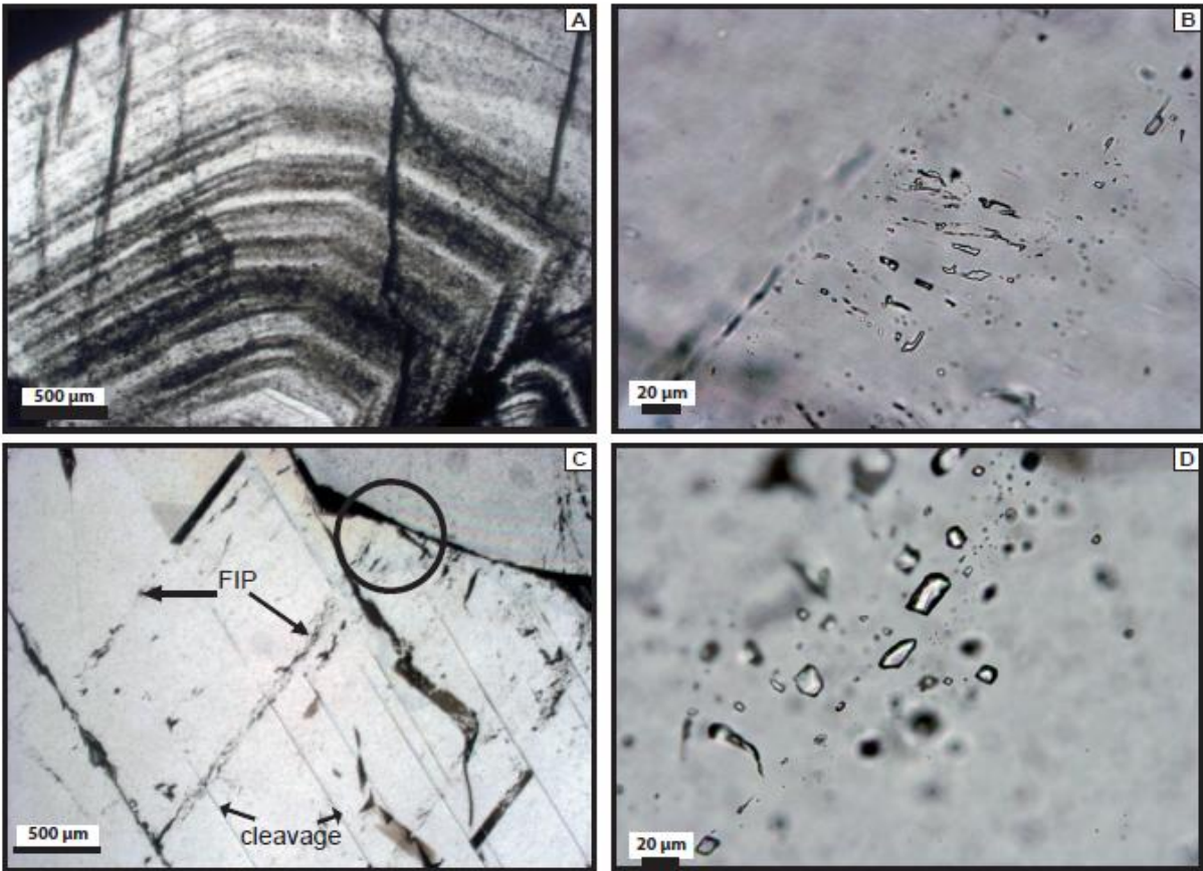


Fig. 4

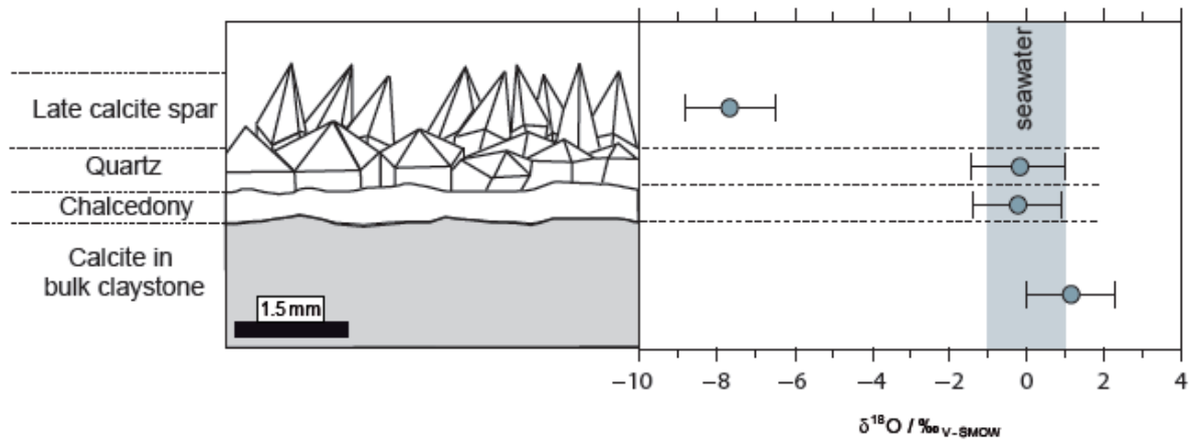


Fig. 5

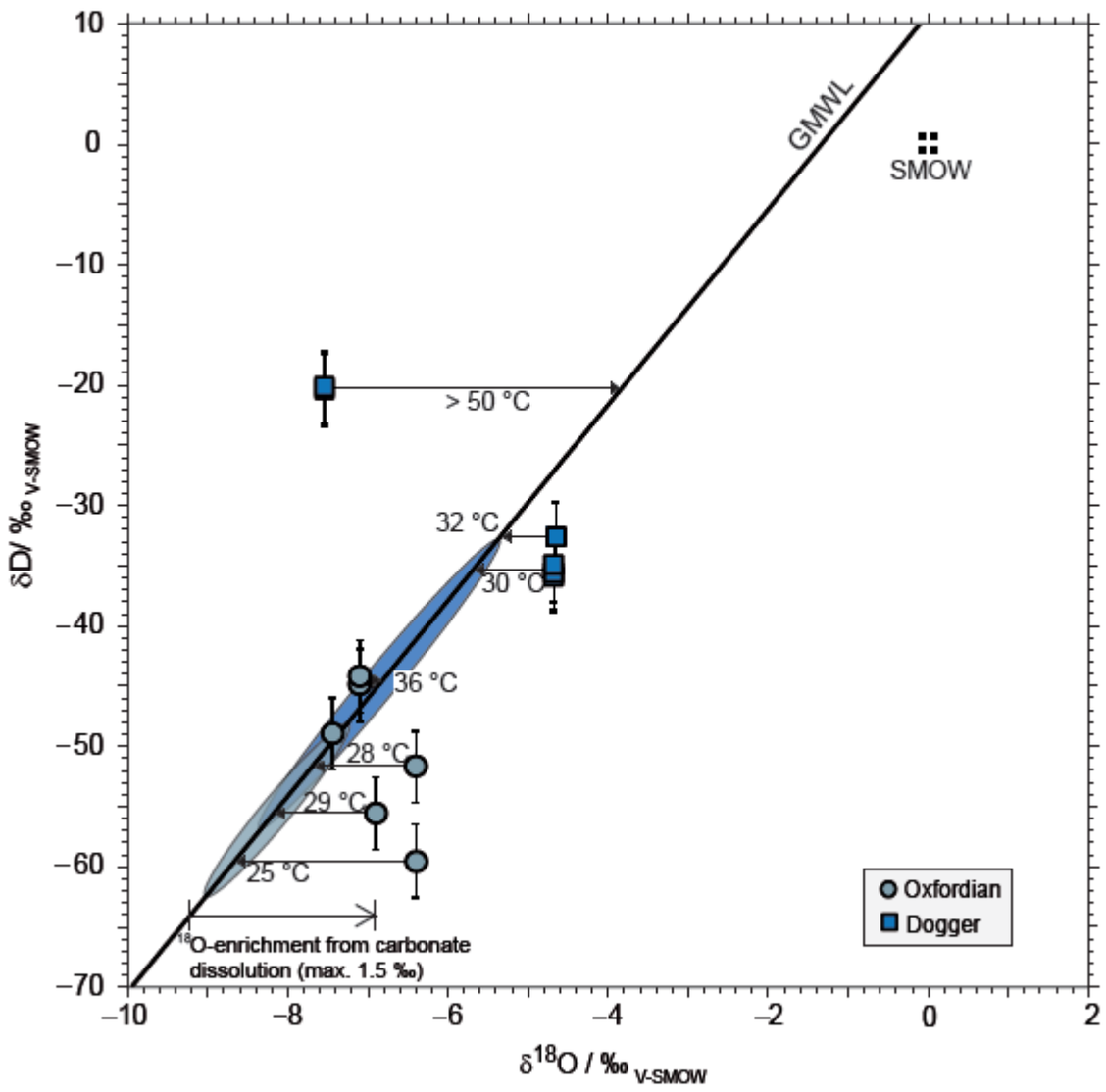


Fig. 6

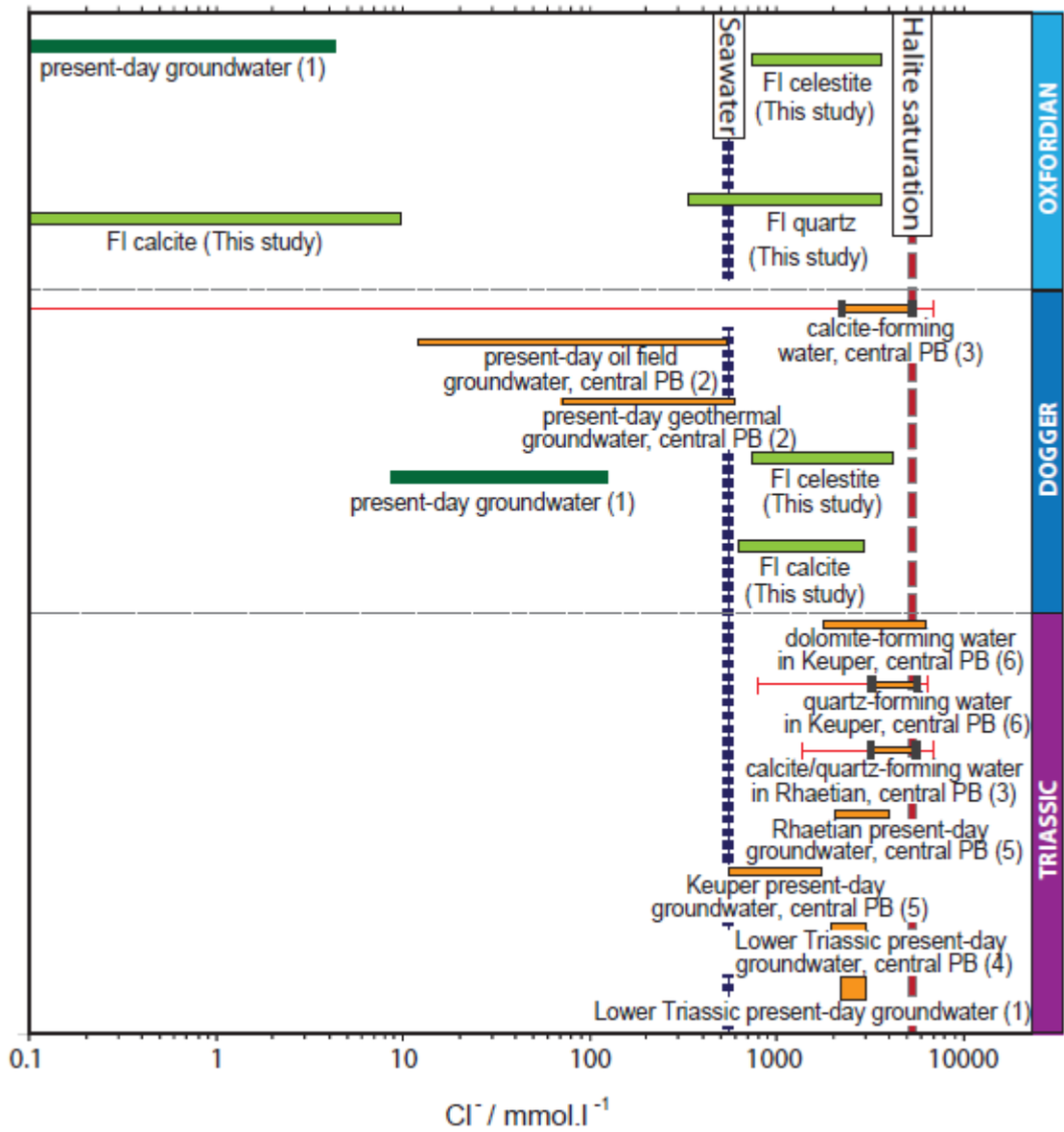


Fig. 7

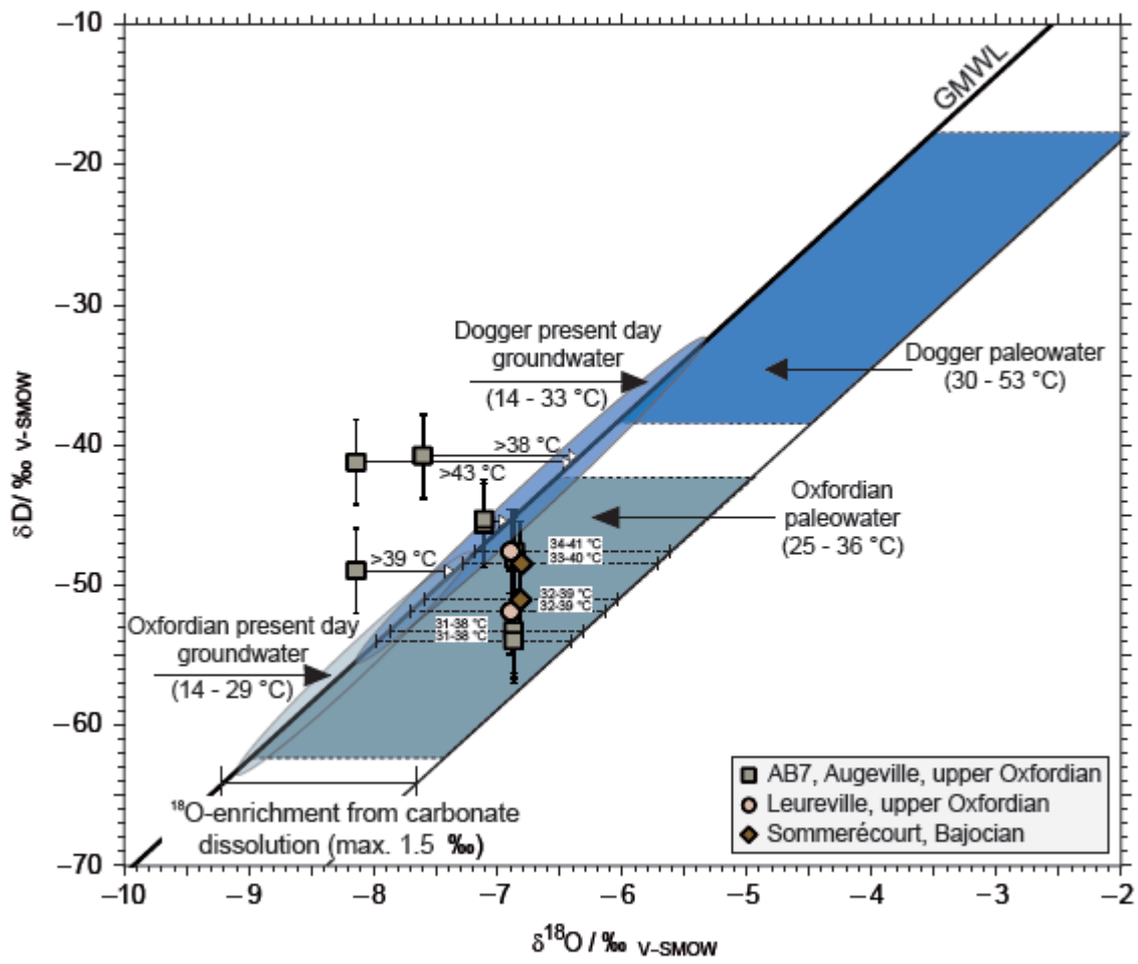


Fig. 8

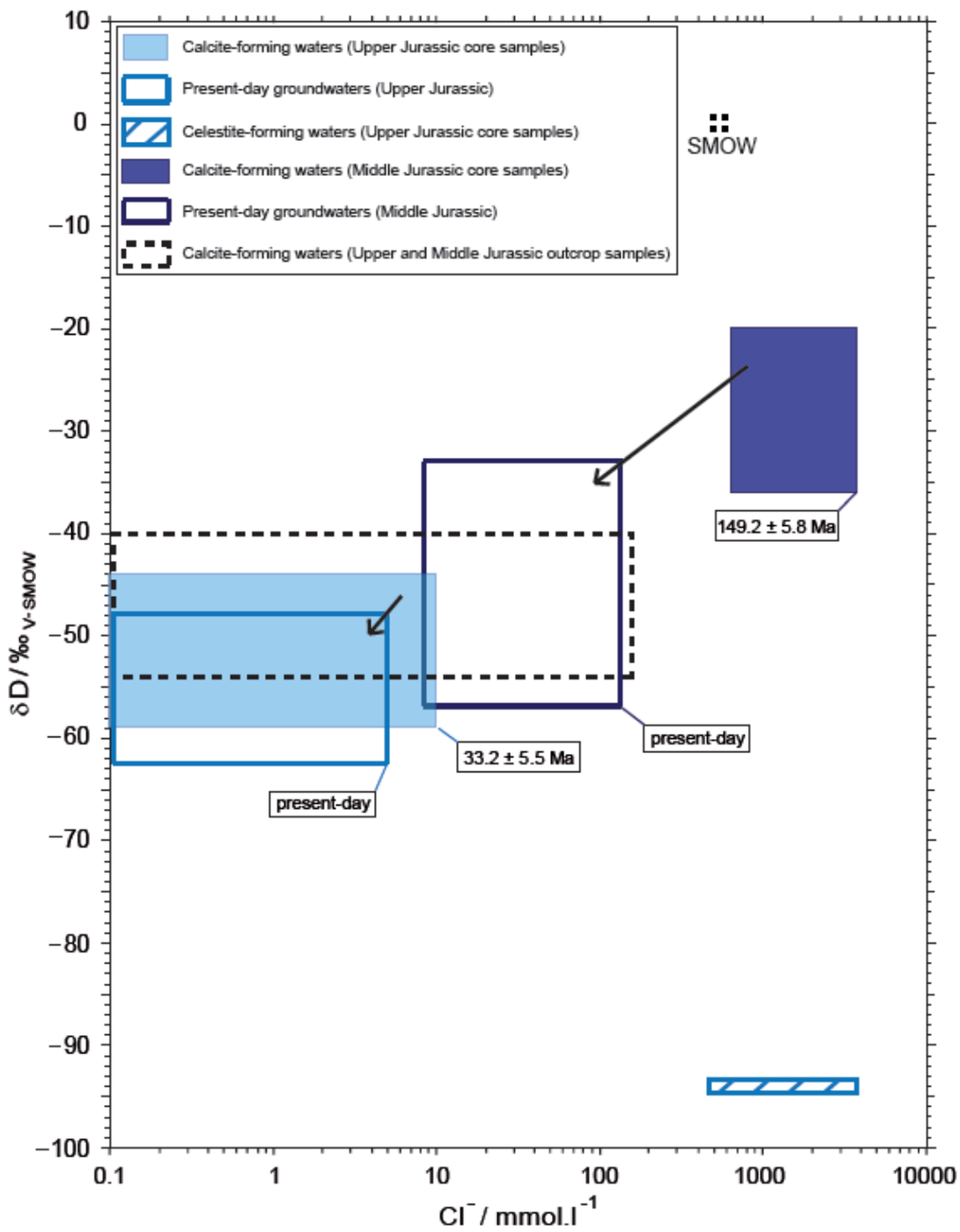


Fig. 9

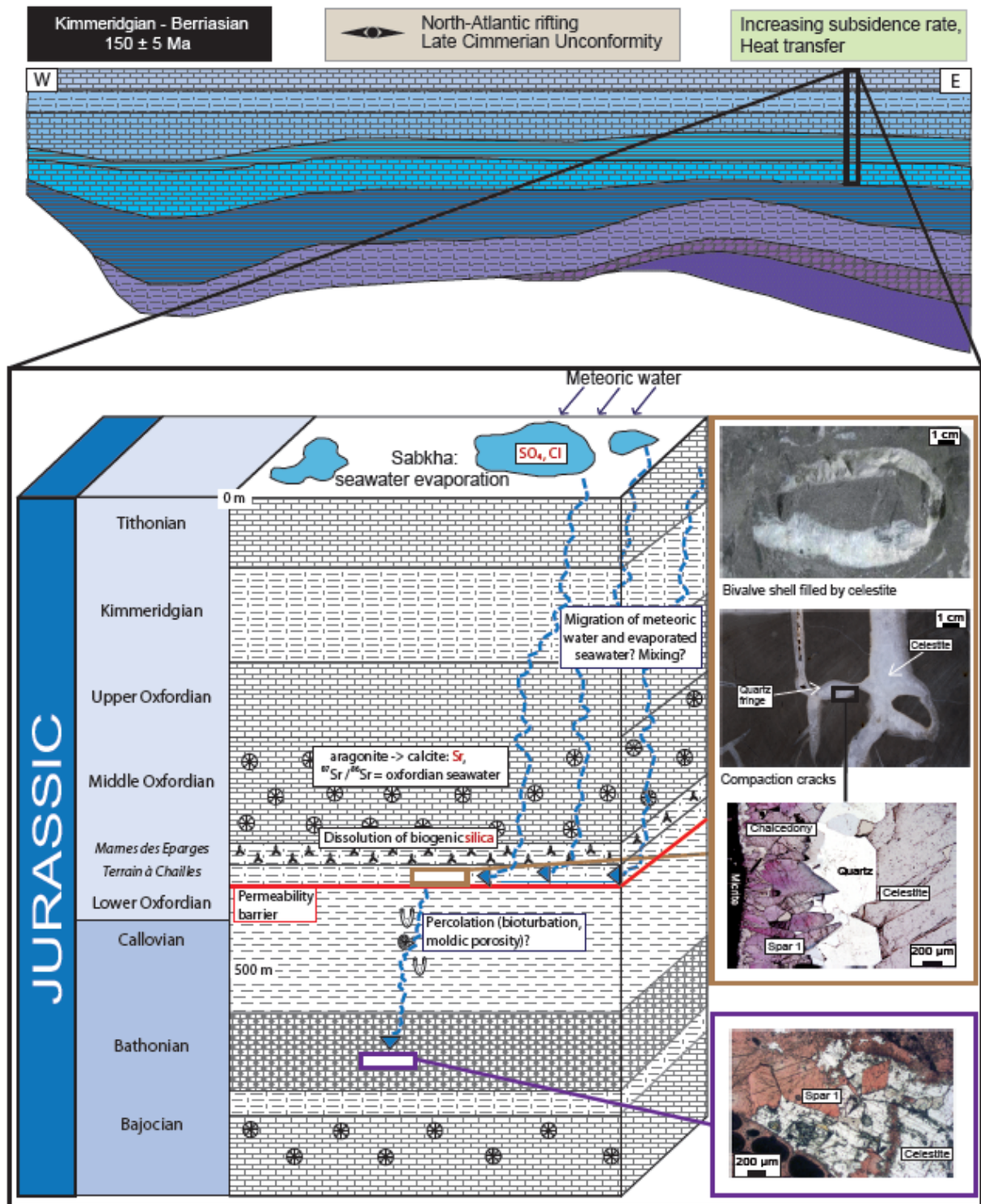


Fig. 10

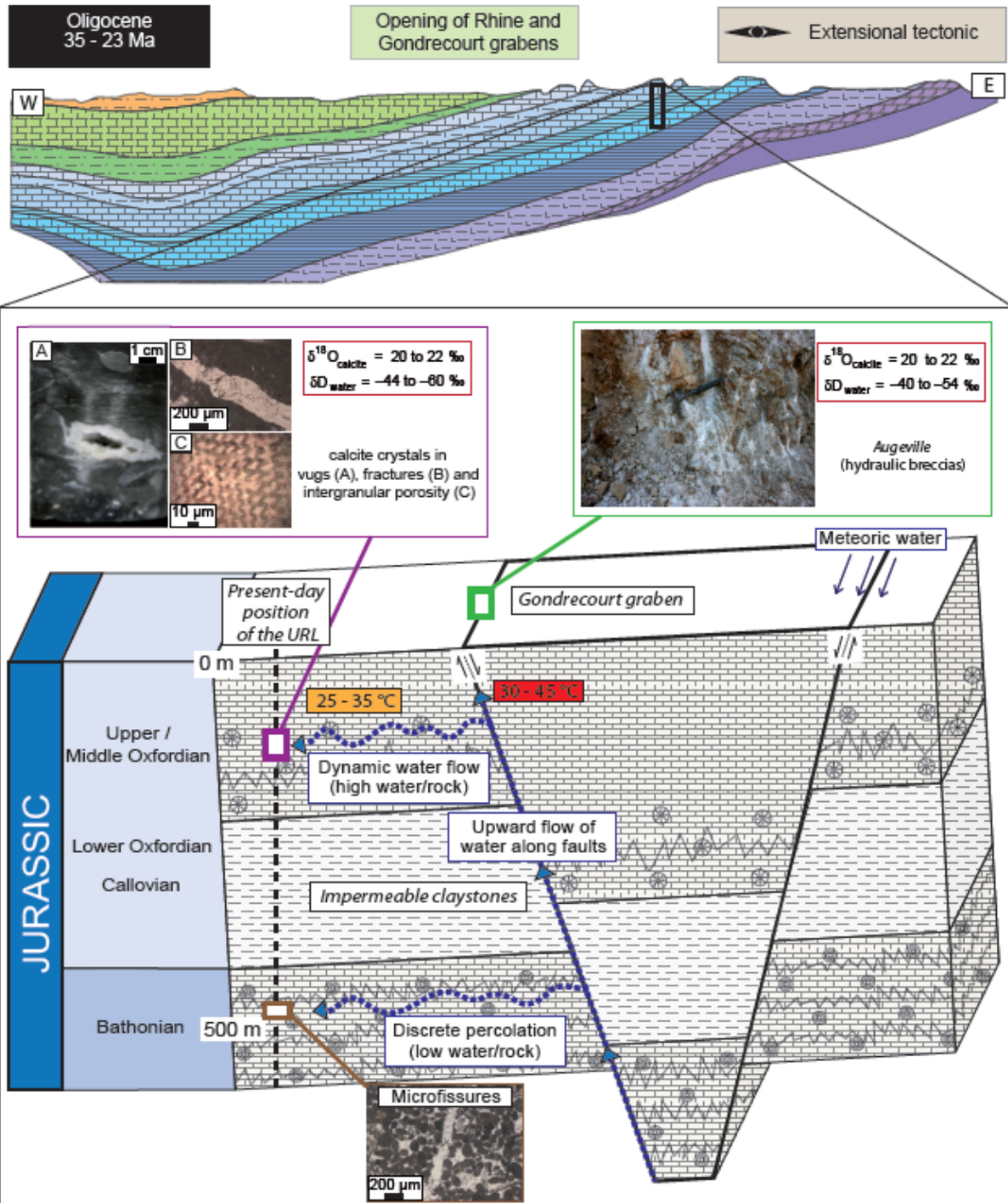


Fig. 11

Table 1

Borehole / Locality	Sample depth - number	Mineral	Stratigraphic position	Sample type	Weight (mg)	H ₂ O (μmol)	δD (‰ v-SMOW)	δ ¹⁸ O _{mineral} (‰ v-SMOW)	δ ¹⁸ O _{H₂O} (‰ v-SMOW) T = 35 °C	δ ¹³ C (‰ PDB)	Chlorinity (mmol l ⁻¹)
PPA	134 m -1	Calcite	Upper Oxfordian	Vug	3213.2	7.06	-59.5	22.4	-6.4	3.0	0 (5)
PPA	134 m -2	id.	Upper Oxfordian	Vug	8211.9	9.14	-51.7	id.	id.	2.9	id.
HTM102	141 m	id.	Upper Oxfordian	Vug	3589.7	19.10	-48.9	21.4	-7.4	3.1	0 (2)
PPA	184 m -1	id.	Upper Oxfordian	Vug	6211.6	33.41	-44.9	21.7	-7.1	2.7	0 (5)
PPA	184 m -2	id.	Upper Oxfordian	Vug	1846.1	32.87	-44.2	id.	id.	2.8	id.
PAX	324 m -1	Celestite	Middle Oxfordian	Compaction crack	1383.5	5.99	-94.0	19.5	/	/	480 - 3640 (8)
PAX	324 m -2	id.	Middle Oxfordian	Compaction crack	/	/	/	19.8	/	/	id.
PAX	340 m	Calcite	Middle Oxfordian	Vug	3606.8	5.74	-55.6	21.9	-6.9	2.9	0 - 50 (8)
EST413	376 m -1	Celestite	Middle Oxfordian	Moldic porosity	1070	2.12	-94.0	19.4	/	/	1300 - 3660 (8)
EST413	376 m -2	id.	Middle Oxfordian	Moldic porosity	/	/	/	19.1	/	/	id.
PAX	380 m	Quartz	Middle Oxfordian	Compaction crack	1747.2	9.98	/	/	/	/	320 - 3620 (21)
EST433	481 m -1	Calcite	Bathonian	Vug	2225.8	98.05	-20.1	21.3	-7.5	0.0	1000 - 1700 (11)
EST433	481 m -2	id.	Bathonian	Vug	dupl.	dupl.	-20.3	id.	id.	0.5	id.
EST433	569 m -1	id.	Bajocian	Vug	2868.2	117.91	-32.7	24.2	-4.7	2.0	650 - 2700 (10)
EST433	569 m -2	Celestite	Bajocian	Vug	/	/	/	/	/	/	760 - 3840 (7)
EST433	660 m -1	Calcite	Bajocian	Vug	2126.6	25.85	-34.9	24.1	-4.7	2.3	unknown (Fe-calcite)
EST433	660 m -2	id.	Bajocian	Vug	dupl.	dupl.	-35.7	id.	id.	2.7	id.
Augeville	AB71	Calcite	Upper Oxfordian	Hydraulic breccia	2020.7	64.16	-48.1	21.9	-6.9	2.5	0 - 40 (6)
Augeville	AB72	id.	Upper Oxfordian	Hydraulic breccia	1080.8	26.50	-47.6	id.	id.	id.	id.
Augeville	AB7-B1	id.	Upper Oxfordian	Hydraulic breccia	2010.4	18.44	-53.6	id.	id.	id.	id.
Augeville	AB7-B2	id.	Upper Oxfordian	Hydraulic breccia	999.9	9.60	-54.0	id.	id.	id.	id.
Augeville	AB7-C1	id.	Upper Oxfordian	Hydraulic breccia	2011.1	37.73	-45.4	21.7	-7.1	2.0	0 (3)
Augeville	AB7-C2	id.	Upper Oxfordian	Hydraulic breccia	1950.3	39.03	-45.7	id.	id.	id.	id.
Augeville	AB7-GC3	id.	Upper Oxfordian	Hydraulic breccia	1012.0	9.43	-40.8	21.2	-7.6	2.0	0 - 30 (4)
Augeville	AB7-GF1	id.	Upper Oxfordian	Hydraulic breccia	2219.7	123.69	-41.2	20.7	-8.2	1.6	0 (2)
Augeville	AB7-GF2	id.	Upper Oxfordian	Hydraulic breccia	1055.0	43.78	-49.0	id.	id.	id.	id.
Leuville	LEU-3	id.	Upper Oxfordian	Hydraulic breccia	1144.1	15.15	-47.6	21.9	-6.9	1.7	0 - 70 (6)
Leuville	LEU-4	id.	Upper Oxfordian	Hydraulic breccia	2005.3	17.99	-51.9	id.	id.	id.	id.
Sommerécourt	SOM2-1	id.	Bajocian	Vug	2131.4	98.47	-48.4	22.0	-6.8	2.9	0 - 160 (3)
Sommerécourt	SOM2-2	id.	Bajocian	Vug	1005.2	28.46	-51.0	id.	id.	id.	id.

Table 2

borehole	Sample depth	Mineral	Sample type	Stratigraphic position	$\delta^{13}\text{C}$ (‰ PDB)	$\delta^{18}\text{O}_{\text{mineral}}$ (‰ V-SMOW)	$\delta^{18}\text{O}_{\text{H}_2\text{O}}$ (‰ V-SMOW) $T = 35^\circ\text{C}$	$^{87}\text{Sr}/^{86}\text{Sr}$
PAX	380 m	Calcite in host rock (marl)	Vug	Lower Oxfordian	2.1	30.0	1.2	/
id.	id.	Chalcedony	id.	id.	/	32.4	-0.3	/
id.	id.	Quartz	id.	id.	/	33.2	-0.2	/
id.	id.	Calcite spar	id.	id.	1.5	21.2	-7.6	/
id.	324 m	Celestite	Compaction crack	id.	/	/	/	0.706950
PPA	360 m	id.	Compaction crack	Middle Oxfordian	/	/	/	0.707545
id.	372 m	id.	Moldic infilling	Lower Oxfordian	/	/	/	0.706924
EST413	376 m	id.	Moldic infilling	Lower Oxfordian	/	/	/	0.707081
EST433	569 m	Celestite	Vug	Bajocian	/	/	/	0.707140

Highlights

- First data combining δD and chlorinity of fluid inclusions in the Jurassic of the Eastern Paris Basin
- Brines are identified in calcite crystallized at around 150 Ma
- Paleofluid temperatures do not exceed 60 °C
- Independent hydrological history in limestone aquifers on both sides of the Callovian-Oxfordian claystones