Research Article

Understanding Fluid Flow during Tectonic Reactivation: An Example from the Flamborough Head Chalk Outcrop (UK)

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1. Introduction

Chalk is a fine-grained carbonate rock constituted mostly of coccolith tests and characterized by a wide range of porosities and low permeabilities. Chalk intervals can act as an effective seal for hydrocarbons in the subsurface [1]. Nevertheless, fractured chalks develop a significant structural porosity and an increased permeability [2, 3]. In this respect, the critical role of fault zones in fluid migration has been demonstrated (e.g., [4, 5]) and the importance of including fault damage zones in reservoir models has been proven [6]. Hence, in order to understand and predict the behavior of faults in subsurface chalk successions, outcrop analogue studies need to be carried out.

The Coniacian-Campanian Chalk of Flamborough Head (UK), located on the southern side of the headland, displays a very complex fracture network, with multiple fracture orientations. The whole chalk section at the study location is intensely fractured by bed-confined normal faults of decimeter to plurimeters length. This fracture network is part of a multiscale polygonal fault system confined to the Coniacian-Campanian chalk deposits [7]. These faults will be referred to as chalk confined fault throughout the article. The outcrops are situated at the intersection of two fault systems:
the E-W trending Howardian–Flamborough Belt and the N-S trending Peak Trough Fault systems. The outcrops have been widely studied and a set of normal fault orientations have been recorded (e.g., [8–10]). Their origin still remains poorly understood and was attributed to a complex multiphase tectonic activity of the North Sea area [11]. Faï-Gomord et al. [7] evidenced polygon-shaped fracture networks and argued for a synsedimentary origin of the normal faults confined to Chalk. Polygonal faults display displacements of decimeter-scale and are interpreted as resulting from a single deformation phase in which faults develop to accommodate compactional strain. This study focuses on a basement fault crosscutting the whole stratigraphic succession [12] and cropping out in Selwicks Bay. The fault core is characterized by a 2 to 3 m wide zone of cemented breccia described in great detail by Starmer [13] and Sagi et al. [14]. Over a few dozen meters on both sides of the fault plane, chalk deposits are highly fractured and display a swarm of calcite veins. Sagi et al. [14] suggest that fluid flow in fractured chalk is especially influenced by the protolith, in particular, the presence of interlayered marl horizons. The origin of the fluid precipitating the calcite, however, has not been addressed, and it is therefore unclear whether it originates from the pressure solution of the chalk itself or from an external source.

This paper examines the origin of the fluid and constrains the local paragenetic history based on a petrographical, geochemical, and microthermometric analyses of the veins. Veins can record the physicochemical conditions of the fluids and provide information about the connectivity of fault conduits by inferring fluid migration pathways (e.g., [15, 16]). Because these features are difficult to characterize in the subsurface, the study of outcrop analogues offers an ideal view of the fractures and fault zones and provides information on potential fluid migration in fractured tight chalk. The presence of fault-related calcite-cemented breccia suggests a hydraulic fracturing, associated with overpressured conditions [17]. Fault-related hydraulic breccias are described in chalk, both in Selwicks Bay (UK) [14] and the Omey Quarry (France) [18]. Results discussed in this paper contribute to the understanding of the behavior of fault zones and associated fluid flow patterns in tight chalks.

2. Geological Setting

Flamborough Head is a headland located on the east coast of Yorkshire in the UK. The study area is located in Selwicks Bay (N 54°7′14.07″, W 0°4′52.769″) at the eastern termination of the Howardian–Flamborough Fault Belt. This is a system of deep normal faults formed during the Late Jurassic–Early Cretaceous (Figure 1). The fault network was then reactivated as reverse faults, between Late Cretaceous till Early Cenozoic [19]. The Howardian–Flamborough Fault belt delimits the southeastern part of the Cleveland Basin from the northern part of the Market Weighton Block [19, 20]. A fault network oriented N-S also crosses the study area: the Peak Trough Fault System, which was reactivated as well during the inversion of the Cleveland Basin, during the Cenozoic [21]. The exact timing of the tectonic inversion ranges from the latest Cretaceous till the Neogene and is linked to Alpine orogenic compression from the south and compression related to Atlantic opening to the north. Throughout the Cenozoic, uplift of the basin continued related to the regional tectonic compression, resulting in uplift of the Cleveland Basin and the formation of the Cleveland Anticline [22]. The studied fault outcropping in Selwicks Bay belongs to the Howardian–Flamborough Fault Belt system and is rooted in the basement [12]. The fault thus crosscut the whole stratigraphic section up to the surface (Figure 1).

The Upper Cretaceous chalks of Flamborough Head were deposited as a postrift sediment [19]. The Cleveland Basin is an exhumed potential petroleum system and an onshore analogue of the Southern North Sea Gas Basin. The Cleveland Basin is a Jurassic to early Cretaceous basin, which was inverted during the Cenozoic, at the time of the Alpine Orogeny [24]. It now forms a significant topographic high in the North York Moors. Upper Jurassic syn-rift marine mudstones rich in organic matter (the Kimmeridge Clay Formation) represent the source rocks for most of the region’s hydrocarbons [25]. In the North Sea, the whole sedimentary succession, both clastic and carbonate are exploited to extract hydrocarbons, with reservoirs ranging from Devonian to Eocene ages. The Upper Cretaceous Chalk, however, is relatively insignificant as a productive reservoir within the UK sector. The maximal burial depth of chalk deposits ranges from 0.8 to 1.8 km, depending on the authors [26–28]. Structural and petrographic studies in Flamborough Head suggest that chalk experienced several episodes of fracturing related to the tectonic reactivation of basement faults [15, 23, 29]. Selwicks Bay contains a 20 m displacement normal fault characterized by an approximately 10 m wide fault core of breccia and pluridecimeters-thick crystalline calcite veins [13].

3. Methodology

3.1. Sampling and Petrography. In order to reconstruct the paleofluid evolution, sampling focused on calcite veins, either associated with plurimeters displacement fault damage zones or infilling of centimeter-tick displacement synsedimentary faults. Chalk samples from undeformed host rock were sampled, as well as fragments of chalk from the fault damage zones, in order to allow a petrographic and geochemical comparison. Samples from the protolith are referred to as matrix samples and stratigraphically correspond to the Coniacian-Santonian. Veins crosscutting relationships were first determined in the field, and several samples were taken of each vein generation (Groups I, II, and III). The calcite veins infilling low displacement synsedimentary faults were also sampled. A total of 35 thin sections were studied with standard petrographic methods, employing observations in transmitted light and cold cathodoluminescence microscopy.

3.2. Conventional Geochemistry. Traditional carbon and oxygen stable isotope analyses (δ13C, δ18O) were performed on 47 powdered carbonate samples obtained with a microdrill. Samples were analyzed at the Institute of Geology and Mineralogy of the University of Erlangen-Nürnberg (Germany).
**Figure 1:** Continued.
Powders were reacted with 100% phosphoric acid at 75°C using a Kiel III online carbonate preparation line connected to a ThermoFinnigan 252 mass spectrometer. Isotopic values are reported in the standard δ notation in per mil relative to V-PDB (Vienna Pee Dee Belemnite). Reproducibility was checked by replicate analyses of laboratory standards and is better than ±0.02‰ (1σ) for carbon and ±0.05‰ (1σ) for oxygen isotope.

$^{87}\text{Sr}/^{86}\text{Sr}$ isotope ratio analyses were performed on seven selected samples at the University of Brussels (ULB). Three measurements were performed on the chalk matrix, including both the host rock (2 samples) and chalk fragments from the breccia (1 sample). Four analyses were performed on calcite veins: 1 sample from Group I as well as Group II and 2 samples from Group III. Measurements were performed with a Multicollector Inductively Coupled Plasma Mass Spectrometer (MC-ICP-MS) following the procedure detailed in Snoeck et al. [30]. After measurements, all the raw data were normalized using a standard-sample bracketing method with the recommended value of $^{87}\text{Sr}/^{86}\text{Sr}$ reported for each sample [31]. Strontium concentrations were acquired from small sample fractions (1 to 3 mg) following the procedure described by Weis et al. [31]. Based on repeated digestion and measurement of reference material, the analytical precision (1 SD) of the procedure outlined above is estimated to be below 3%.

A total of 32 representative samples were analyzed for their elemental composition (Mg, Ca, Sr, Fe, Mn, Na, K, Al, Ba, Ni, P, S, Ti, and Zn). Around 50 mg of powdered samples were digested using the so-called 4 acids digestion method, in a Teflon beaker on a hot plate. Samples were then measured using the Inductively Coupled Plasma Optical Emission Spectroscopy (ICP-OES) Varian 720ES, Varian Inc., Walnut Creek, CA, USA. Analytical error is less than 5%.

3.3. Fluid Inclusion Microthermometry. Double polished 150 µm thick wafers were studied under transmitted and UV-fluorescent light microscopy to select inclusions suitable for microthermometry, following the procedures described in detail by Goldstein and Reynolds [32] and Hurai et al. [33]. The method used for the preparation of the aforementioned double polished sections is described in Muchez et al. [34] and has been proven successful for low-temperature carbonates in a diagenetic system. In order to overcome metastability of the monophase fluid inclusions, samples were put in a freezer at −20°C. This procedure stimulates the formation of vapor bubbles in metastable inclusions without freezing of the samples (e.g., [32]). Measurements were done on a Linkam MDS-600 heating-cooling stage mounted on an Olympus BX51 microscope. The stage was calibrated by synthetic Syn Flinc™ fluid inclusion standards. Two-phase aqueous inclusions were heated before freezing in order to obtain the total homogenization temperatures (Th). Monophase inclusions were artificially stretched by repeated heating of the sample up to 200°C and subsequent freezing to −100°C. These successive cycles caused the formation of larger vapor bubbles. Vapor bubbles which were created by artificial stretching have not been used to determine homogenization temperatures but only for the system (composition, e.g., H$_2$O-NaCl or H$_2$O-CaCl$_2$) and the salinity [35]. Subsequent freezing/heating cycles were applied to acquire the first and final ice melting temperatures. All melting temperatures were
recorded in the presence of a vapor bubble, in order to avoid metastable melting [36]. Due to difficulties with the observations related to small inclusion sizes, final melting temperatures of ice have been acquired by sequential freezing [37]. To accurately measure the first and final ice melting temperatures of the fluid inclusions, the stage was heated at a very slow rate (0.2°C per minute). In carbonate rocks, it is common to observe stretching of the inclusions related to the freezing process. Salinities are reported in equivalent weight percent NaCl (eq. wt% NaCl) and were calculated from the temperature of ice melting using the equation of Bodnar [38]. Calculation of the isochors of the aqueous fluid inclusions was carried out using the equation of state of Zhang and Frantz [39].

3.4. Carbonate Clumped Isotope (Δ47) Measurements and Data Processing. The Δ47 thermometry is based on the temperature-dependent preference of 13C and 18O isotopes to bond with each other within the mineral lattice at low temperatures. Carbonate clumped isotopes measurements were made at the Institut de Physique du Globe de Paris (IPGP). A detailed description of the analytical setup is given in Bonifacie et al. [40] and is only briefly summarized hereafter. For each analysis, about 5 mg of powdered calcites were digested for 20 minutes in 104% phosphoric acid at 90°C. After digestion, the sample CO2 was stripped from trace amounts of water and sulfur or organic compounds before being introduced into the mass spectrometer. The mass spectrometer is a Thermo Fisher MAT 253 operated in dual-inlet mode and configured to simultaneously measure masses 44 through 49. Carbonate clumped isotopes were simultaneously measured with multiple analyses of different CO2 gases (with δ13C and δ18O values covering the range of the unknown samples) that were brought to thermodynamic equilibrium at either 1000°C or 25°C and analyzed interspersed with carbonate standards and unknown samples. As our samples were digested at 90°C, an acid correction factor of 0.082‰ (following [42]) was added in order to report the data in a 25°C acid digestion frame to follow the common use. All samples were analyzed three times in three discrete analytical sessions separated by several weeks (i.e., with ETF correction frames constructed with various sets of data on CO2 standards equilibrated at known temperatures) in order to eliminate possible bias from potential error in constructing one of the three discrete correction frames. To ensure accuracy of Δ47 data and the entire data reduction process described above, we routinely analyzed two carbonate reference materials (Carrara marble and 102-GCAZ01b, also reported by Dennis et al. [41] and many other studies) distributed along the unknown samples in all runs. The standard deviations of carbonate reference materials are typically of ±0.014‰, consistent with long-term external reproducibility for Δ47 measurements at IPGP (1 SD, n > 300; [40, 43]). Traditional δ18O and δ13C data were acquired as part of each Δ47 analyses and 17O corrections were made using the 17O parameters from Santrock et al. [44]. In order to account for the temperature dependence of oxygen isotope fractionation between CO2 gas and calcite resulting from the reaction with phosphoric acid at 90°C in a common acid bath, a fractionation factor of 1.00811 was used [43].

The Δ47 data were converted into temperatures using the interlaboratory composite Δ47-T calibration recently published for all carbonate minerals with crystallization temperatures ranging from −1°C to 300°C [40]. Because any Δ47 measurement comes with simultaneous determination of the oxygen isotopic composition of the analyzed carbonate (δ18O_carb), Δ47 thermometry provides independent estimates of the carbonate formation temperature and of the oxygen isotopic composition of the mineralizing fluid (δ18O_water). Δ47 measurements at IPGP were carried out using the equation of state of Zhang and Frantz [39].

4. Results

4.1. Petrography

4.1.1. Macroscopic Observations. A normal fault was studied at Selwicks Bay, striking ENE-WSW, and dipping steeply (N70°) to the NNW. The fault zone is characterized by calcite cementation and presence of cemented brecciated zones [13, 14]. The main fault core forms a 5 m wide promontory on the cliffs. The damage zone of the fault is characterized by a much higher density of calcite veins compared to the surrounding wall rocks. The intensity of calcite veining reduces progressively when moving further away from the fault core zone. In the core zone of the main fault at Selwicks Bay, field observations evidenced three generations of calcite veins; namely,

(i) Group I veins correspond to thin crack and seal like veins, characterized by a network of mostly parallel thin veins measuring a few micro- to a few millimeters wide. They are oriented perpendicular to stratigraphy. In some places, the stratigraphy is tilted, and veins of Group I are oriented perpendicular to the stratigraphy, indicating that veins occurred prior to the tilting. Group I veins usually crosscut compaction stylolites (Figure 2(a)) but are sometimes crosscut by the latter (Figure 2(b));

(ii) Group II veins consist of a breccia with angular rock fragments healed by calcite cement. Clasts are composed of chalk and occasionally chert with sizes ranging from millimeter to decimeter (Figures 2(c)–2(e)). Both the angular shape of the fragments...
Figure 2: Macroscopic observations of the calcite veins. (a) Group I vein generation characterized by a network of pseudo parallel thin veins. (b) A vein from Group I crosscut and displaced by a compaction stylolite. (c) Hydraulic breccia and its typical jigsaw texture. (d) Hydraulic breccia crosscutting the stratigraphic section and compaction stylolites. (e) Hydraulic breccia (with Group II veins) intruded by Group III vein (bright white color). (f) Group III vein crosscutting compactional stylolites. (g) Large calcite crystals infilling porosity along the main fault plane (Group III veins).
and the jigsaw puzzle configuration reflect a limited transport, in conjunction with fast crystallization pointing towards a hydraulic type of fracturing. Chalk fragments include Group I veins, constraining the relative chronology of the vein generations;

(iii) Group III veins are to the last vein generation identified. Veins III crosscut the hydraulic breccia. They are characterized by a bright white color and large calcite crystals. In the main brecciated zone, calcite crystals up to tens of centimeters in width fill up the macroporosity in the vicinity of the fault plane (Figures 2(f) and 2(g)).

Within the Flamborough Head area, on both sides of Selwicks Bay fault, decimeter to few meters long bed-confined normal faults are present. They are part of the polygonal fault systems described by Fay-Gomord et al. [7], which include fault systems of different scales from centimetric to decametric. Some of those faults are cemented. Calcite cementation is observed in the south of South Landing and north of Thornwick areas, up to 1 km away from Selwicks Bay fault. These veins were also sampled and referred to as chalk confined fault (CCF) veins. Some of these faults were reactivated as tectonic stylolites, confirming the crosscutting relationships established during field observations. Group I veins consist of calcite crystals, usually ranging in size between 10 and 50 \( \mu \text{m} \) (Figures 3(a)–3(c)). They display mutual crosscutting relationships with bedding-parallel stylolites. The hydraulic breccia cement (Group II veins) is characterized by sparite calcite crystals measuring usually several 100 \( \mu \text{m} \). Cement commonly displays a drusy mosaic texture (Figure 4(b)), whereby crystal size increases from the border of the chalk fragments towards the centers of the veins. Impurity fringes are commonly observed along chalk fractures in the hydraulic breccia and also contain rhombohedral dolomite crystal ghosts (Figures 3(b) and 3(c)). Dolomite ghosts measure 50 to 150 \( \mu \text{m} \), which still show no vapor bubble development. The monophasic aqueous inclusions which still show no vapor bubble development are two-phase aqueous inclusions, which most inclusions are monophase aqueous inclusions. Few fluid inclusions are two-phase aqueous inclusions, which range in size from \(-8\) to \(12 \mu \text{m} \), with a relatively constant liquid: vapor ratio. The monophase liquid aqueous inclusions range from a few microns to \(\sim 10 \mu \text{m} \) and are interpreted to have been entrapped below \(\sim 50^\circ\text{C} \) (cf. [32]). Monophase inclusions which still show no vapor bubble development after cooling up to \(-20^\circ\text{C} \) in a freezer are most likely not metastable (Muchez & Slobodnick, 1996).

4.2.2. Strontium Isotopes. The \(^{87}\text{Sr}/^{86}\text{Sr} \) ratios in matrix chalk samples range from 0.7074 to 0.7075. The results are in good agreement with the Upper Cretaceous marine carbonate signal (0.7073–0.7076) reported by Mearon et al. [46]. The \(^{87}\text{Sr}/^{86}\text{Sr} \) ratios obtained in veins display values significantly higher. \(^{87}\text{Sr}/^{86}\text{Sr} \) equals 0.7103 in the sample of Group I vein generation, 0.7085 in the cement of the hydraulic breccia (Group II), and 0.7103 in the last vein generation (Group III).

4.2.3. Trace Elements. The elemental concentrations are shown in Table 1. The chemical compositions of the host rock and the veins display very distinct signatures. Veins, independently of their generation (e.g., Groups I, II, and III), exhibit low concentrations of trace elements. Chalk matrix samples generally show higher Al, Fe, and K content than the calcite veins, while no significant differences in Ca and Mg concentrations are observed. The Al and K contents relate to the presence of noncarbonate constituents, especially clays in the chalk. The high Fe and S concentration of the chalk may be associated with the presence of some pyrite in the matrix. Fe concentration equals 491 ppm on average for matrix samples. Veins are characterized by significantly lower Fe concentrations, with Group II and III veins displaying the lowest Fe concentration (110 ppm on average) and Group I veins with higher values (232 ppm on average). The geochemical signature of the CCF veins, infilling the polygonal fault systems is very similar to Group I and II veins.

4.3. Fluid Inclusion Microthermometry. Numerous aqueous fluid inclusions displaying various degree of deformation are identified along the crystal growth planes (Figure 6). The fluid inclusions selected for measurements are primary in origin and show no indication of stretching, necking down, or deformation. Optical and fluorescence microscopy revealed that most inclusions are monophase aqueous inclusions. Few fluid inclusions are two-phase aqueous inclusions, which range in size from \(-8\) to \(12 \mu \text{m} \), with a relatively constant liquid: vapor ratio. The monophase liquid aqueous inclusions range from a few microns to \(\sim 10 \mu \text{m} \) and are interpreted to have been entrapped below \(\sim 50^\circ\text{C} \) (cf. [32]). Monophase inclusions which still show no vapor bubble development after cooling up to \(-20^\circ\text{C} \) in a freezer are most likely not metastable (Muchez & Slobodnick, 1996).

Difficulties with the measurement of the first melt temperature are explained by the limited amount of hydrohalite and ice that melt at the eutectic temperature (Te) of \(\sim 21.2^\circ\text{C} \), in combination with a small inclusion size. The first melt related observations made below Te is the appearance of rounded ice crystals with a fluid in between them. Freezing
behavior shows additional evidence for a relatively low saline H2O-NaCl system. The temperature of metastable freezing (Tfr) is observed by an instantaneous volume decrease, deformation, or jump in bubble position. This "jerk" or "collapse" of the vapor bubble, in the observed temperature range of −34°C to −50°C, is typical behavior for H2O-rich inclusions (Roedder, 1984). The aforementioned range in Tfr measurements can be explained by salinity and inclusion size variations [47].

Group I veins do not contain any fluid inclusions large enough for microthermometric analyses.

Group II veins (e.g., hydraulic breccia cement) are characterized by homogenization temperatures ranging from 34 to 48°C. The temperature of first melting is difficult to observe and the lowest values are below −10°C. The final melting temperatures range from −7.2 to −0.1°C. Using the equation of Bodnar [38] and assuming a H2O-NaCl system based on the freezing temperatures (Tfr), the final melting temperatures correspond to a salinity ranging from 0.3 to 10.7 eq. wt% NaCl.

The homogenization temperatures of inclusions in Group III veins are between 37 and 53°C. The temperature of first
melting is difficult to observe and is below $-7^\circ$C. Within a H$_2$O-NaCl system, based on the final melting temperatures, ranging from $-4.6$ to $-0.1^\circ$C, the calculated salinities range from 0.2 to 7.3 eq. wt% NaCl.

All vein generations display similar homogenization temperatures and a range of salinity values. Most salinity values are below 1 eq. wt% NaCl and reflect a meteoric water signal, as shown in Figure 6(c). The spread of values between a meteoric signal and 10 eq. wt% NaCl could be due to the mixing of meteoric water with a saline fluid or to the dissolution of evaporites.

The average homogenization temperature measured is $45^\circ$C. Using a temperature of $45^\circ$C and two different salinity values of, respectively, 7.7 and 0.4 eq. wt% NaCl, the isochores of the latter two fluids have been constructed in a P-T diagram (Figure 6(d)). The burial depth is estimated around 1 km since veins crosscut compaction stylolites and are crosscut by compaction stylolites, formed from a depth of $\sim 800$ m in chalk [48]. Using the appropriate isochores (lines of equal density) and a typical hydrostatic pressure of 10 MPa/km, calculated trapping temperatures are 52 and 53$^\circ$C. Using a typical lithostatic pressure gradient of 25 MPa/km for the same samples, calculated temperatures are 60 and 62$^\circ$C.

4.4. Temperature and $\delta^{18}O_{water}$ from $\Delta_{47}$ Data. Data obtained from clumped isotope measurements on three carbonate groups are reported in Table 2. For one analyzed sample, the three $\Delta_{47}$ measurements are showing external

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**Figure 4:** Microscopic observations under plane-polarized transmitted light (1) and cathodoluminescence (2). (a) Group I veins with visible sector zonations. (b) Hydraulic breccia cemented by drusy calcite cement. (c) Group I veins with sector zonations visible under cathodoluminescence.
reproductibilities that are consistent with the ±0.014‰ long-term external reproducibility obtained on homogeneous carbonate standards over several years of analyses. This attests to the homogeneity of the samples and the good quality of the isotopic measurements, with three groups showing an average Δ47 value of 0.588 ± 0.012‰ (n = 3 samples and 9 measurements). The latter suggests indistinguishable formation temperature (average temperature of 60 ± 5°C, 1 SD), for the three samples from each vein generation. The calculated δ18Owater values of the three samples are also indistinguishable within uncertainty, with average values of 0.5 ± 0.9‰ (1 SD).

5. Discussion

5.1. Origin of the Fluid. Several pulses of fluid flow were identified based on the crosscutting relationships, as confirmed by the petrographical analyses. Nevertheless, similar trace element and isotopic compositions of the different vein generations reveal one common fluid source. The strontium isotopic composition of the veins possesses a radiogenic signal, which does not match that of the Upper Cretaceous seawater. The high 87Sr/86Sr ratio is probably related to the interaction of the fluid with minerals containing radiogenic 87Sr. Enrichment in 87Sr may occur when the mineralizing fluid had been in contact with siliciclastic aquifers. Trace element analyses, which revealed a low concentration of Fe or Mn in the calcite veins, suggest that the aquifers were probably composed of sandstones. The spread of salinity values measured by microthermometry analyses supports a dominant meteoric origin of the fluid, which has either been mixed with saline fluids or enriched in salt due to the dissolution of evaporitic minerals. In the Cleveland Basin, the Sherwood Sandstone Group represents an aquifer, up to 450 m thick [49], which belongs to the Permo-Triassic sandstones group, the second most important aquifer in the UK [50]. If the infiltration area of the Sherwood Sandstone aquifer is located hundreds of kilometres away, the Sherwood Sandstone Formation is still present just 1 km below the chalk in the study area (Figure 1). The Sherwood Sandstone Group forms the most important aquifer in Northern England and the meteoric water migrated in the formation far away from the recharge area as reported in several papers. A number of large towns obtain their water supplies at least partly from the Permo-Triassic sandstones, among them are Manchester, Liverpool, Birmingham, Leeds, Doncaster, and Nottingham [51]. The sedimentology and diagenesis of the Sherwood sandstones have been widely studied: Burley [52] addressed the composition of the sandstone reservoir which contains several Sr-bearing minerals, including plagioclase feldspars, as well as igneous and metamorphic rock fragments. Worden et al. [53] described highly saline formation waters within the Triassic Sherwood Sandstone of the Wessex Basin and suggested that a meteoric fluid became enriched in salt due to the dissolution of evaporites. The lithostratigraphy of the Cleveland Basin reveals the presence of halite deposits, both underneath the Permo-Triassic sandstones and interbedded within the Upper Triassic Mercia Mudstone Group, which overly and confine the Sherwood sandstones (Figure 1(b)). The latter is considered the most probable source of salt within the Sherwood Sandstone aquifer [53].

Fluid inclusion microthermometric analyses display salinity values supporting dominantly meteoric water. Nevertheless, the δ18Owater value of the mineralizing fluid calculated from clumped isotope measurements range from −0.2 to 1‰, while the meteoric water signature usually ranged from −3 to −10‰ (SMOW) at the corresponding latitude from the Permian to Late Cretaceous [54]. If the fluid remained stored in an aquifer for sufficiently long time period, it is probable that δ18Owater reflects an extended water-rock interaction under low fluid to rock ratio [55]. In addition, some samples display low δ13C values and two samples especially possess negative values, which probably reflect interference from a depleted source.

The diagenesis of the Sherwood Sandstone has been described from several locations throughout the UK, the Irish Sea, and the North Sea (e.g., [52, 56, 57]). Deep burial diagenetic processes led to the precipitation of different cement phases, including ankerite and anhydrite cements, which represent several percentages of the total bulk rock volume, occasionally reaching up to 17% [54, 58, 59]. These cements, which overgrow previous mineral phases, reportedly yield original δ18Owater (SMOW) values ranging from 0 to 5‰ (2‰ on average) [54]. Ankerite precipitation in the Sherwood Sandstone probably led to iron depletion in the formation water of the Sherwood Sandstone. The water could have been remobilized during tectonic reactivation giving rise to the cementation of calcite veins within the Chalk. This hypothesis could also explain the low iron concentrations measured within the calcite veins (200 ppm on average), which also yield a radiogenic signal.
Table 1: Trace elements concentrations measured by ICP-OES.

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<td>793</td>
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<td>7</td>
<td>37.3</td>
<td>1761</td>
<td>15</td>
<td>663</td>
<td>468</td>
<td>1813</td>
<td>436</td>
<td>15</td>
<td>199</td>
<td>151</td>
<td>1028</td>
<td>80</td>
<td>15</td>
</tr>
</tbody>
</table>

| CCF veins | 37 | 39 | 72 | 44 | 213 | 30 | 1350 | 413 | 1 | 12 | 44 | 1229 | 3 | 2 |
| 8     | 38.9| 87  | 5   | 75  | 31  | 1457| 312 | 2   | 18  | 37  | 1147 | 4 | 2  |
| 9     | 38.8| 45  | 1   | 770 | 17  | 1433| 327 | 2   | 6   | 29  | 777  | 1 | 2  |
| 10    | 38.6| 115 | 6   | 304 | 48  | 1430| 413 | 1   | 14  | 32  | 1907 | 7 | 4  |
| 11    | 38.6| 83  | 3   | 26  | 31  | 1697| 404 | -   | 19  | 23  | 1972 | 3 | 2  |
| 12    | 38.9| 62  | 5   | 331 | 32  | 1408| 404 | 0   | 16  | 33  | 1613 | 2 | 2  |
| 13    | 38.6| 112 | 7   | 47  | 46  | 1307| 424 | 1   | 15  | 42  | 1597 | 4 | 2  |
| 14    | 38.7| 43  | 322 | 37  | 22  | 951 | 496 | 0   | 2   | 118 | 373  | 1 | 2  |
| 15    | 38.8| 29  | 3   | 114 | 16  | 1115| 522 | 0   | 3   | 40  | 447  | 1 | 1  |

| Group I veins | 39 | 275 | 5 | 233 | 106 | 1504 | 402 | 35 | 48 | 159 | 1309 | 11 | 4 |
| 16    | 38.6| 185 | 3   | 125 | 60  | 1304| 433 | -  | 63  | 113 | 951  | 6 | 1  |
| 17    | 41.7| 144 | 4   | 389 | 96  | 1614| 428 | 19  | 21  | 236 | 1598 | 7 | 1  |
| 18    | 39.0| 38  | 2   | 59  | 18  | 1389| 283 | 21  | 9   | 17  | 958  | 1 | 1  |
| 19    | 38.1| 355 | 9   | 370 | 134 | 1655| 426 | 125 | 58  | 145 | 1602 | 11| 6  |
| 20    | 38.0| 346 | 6   | 277 | 137 | 1488| 431 | 4   | 67  | 359 | 1151 | 13| 5  |
| 21    | 38.3| 584 | 5   | 176 | 190 | 1571| 410 | 4   | 70  | 84  | 1595 | 28| 4  |

| Group II veins | 39 | 229 | 11 | 100 | 78  | 1540 | 494 | 10 | 57 | 86  | 1349 | 11 | - |
| 22    | 38.6| 146 | 5   | 53  | 68  | 1435| 429 | 17  | 39  | 115 | 1667 | 7 | - |
| 23    | 39.3| 312 | 17  | 146 | 88  | 1644| 559 | 3   | 74  | 56  | 1030 | 14| - |

| Group III veins | 39 | 52  | 6  | 79  | 31  | 1601| 444 | 13 | 17 | 42  | 1495 | 8 | 3 |
| 24    | 38.9| 49  | 5   | -   | 57  | 1609| 473 | 29 | -   | 54  | 1474 | 3 | - |
| 25    | 38.9| 56  | 6   | 35  | 33  | 1567| 471 | 30 | 20  | 85  | 1415 | 3 | - |
| 26    | 38.6| -   | 2   | 41  | 19  | 1484| 436 | -  | -   | -   | 1429 | - | - |
| 27    | 39.0| -   | 2   | 341 | -   | 1435| 462 | 15 | -   | -   | 944  | 49| - |
| 28    | 38.8| 35  | 6   | 16  | 18  | 1753| 449 | 8   | 18  | 31  | 1702 | 1| 3  |
| 29    | 38.5| 28  | 6   | 23  | 29  | 1751| 447 | 12  | 18  | 24  | 1698 | 1| 6  |
| 30    | 37.1| 58  | 6   | 125 | 29  | 1572| 437 | 0   | 9   | 33  | 1151 | 5| 2  |
| 31    | 40.1| 66  | 17  | 26  | 34  | 1679| 413 | -  | 25  | 39  | 2357 | 2| 1  |
| 32    | 38.6| 69  | 2   | 28  | 27  | 1555| 410 | 0   | 10  | 27  | 1286 | 2| 2  |

CCF: chalk-confined fractures, ‘-’: values below detection limit; LOQ: limit of quantification.
5.2. Relative Timing of Events and Evolution of Fluid Flow. The diagenetic processes affecting the Flamborough Chalk deposits are summarized in Figure 8 and are shown in relation to the regional burial history diagram proposed by Emery [23].

Early Faulting and Fluid Flow Event. The petrographical study of the breccia revealed a large number of impurities and rhombohedral crystal ghosts, concentrated in clusters along one side of some chalk fragments. Crosscutting relationships with calcite sparitic cement (Vein Group II) show that the impurities-rich fringes predate vein cementation. The presence of rhombohedral ghosts suggests the formation of dolomite rhombs prior to the breccia cement mineralization. A possible source of Mg necessary for dolomite precipitation could be related to smectite-illite recrystallization processes. Smectite minerals are converted to ordered smectite-illites as temperatures increase with burial depth, releasing free ions in solution. In the Cleveland Basin, the Chalk Group is underlain by a hundred meters thick succession of clays and clay-rich deposits. Previous studies demonstrated that the Lower Jurassic Whitby Mudstone Formation [60], the Lower Cretaceous Speeton Clays [61, 62], or the Upper Jurassic Oxford and Kimmeridge clays [63, 64] contain illite-smectite (I-S) mixed layer clays with up to 90% illite. The overpressure created by the smectite to illite conversion in the Cleveland Basin may have led to fluid migration into the overlying lithologies [65], including the Chalk. We suggest Mg$^{2+}$ enriched fluids migrated along the main fault zone and within the preexisting connected fault system of the chalk, resulting in the crystallization of dolomite rhombs in veins along fault planes. The migration of this fluid predates the
Table 2: Data from clumped isotopes analyses.

<table>
<thead>
<tr>
<th>Sample Group</th>
<th>$\delta^{13}$C (‰, VPDB)</th>
<th>$\delta^{18}$O (‰, VPDB)</th>
<th>Δ47 ‰, CDES</th>
<th>Av. Δ47 ‰, CDES</th>
<th>±1 S.D. (‰)</th>
<th>T. (°C)</th>
<th>$\delta^{18}$Owater (SMOW)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample Group I</td>
<td>2.54</td>
<td>-7.74</td>
<td>0.582</td>
<td></td>
<td></td>
<td>60</td>
<td>± 3</td>
</tr>
<tr>
<td></td>
<td>2.56</td>
<td>-7.74</td>
<td>0.581</td>
<td>0.588</td>
<td>0.012</td>
<td>1.0</td>
<td>± 0.8</td>
</tr>
<tr>
<td></td>
<td>2.54</td>
<td>-7.69</td>
<td>0.602</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sample Group II</td>
<td>2.61</td>
<td>-8.21</td>
<td>0.595</td>
<td></td>
<td></td>
<td>56</td>
<td>± 3</td>
</tr>
<tr>
<td></td>
<td>2.47</td>
<td>-8.19</td>
<td>0.601</td>
<td>0.597</td>
<td>0.004</td>
<td>-0.1</td>
<td>± 0.2</td>
</tr>
<tr>
<td></td>
<td>2.60</td>
<td>-8.25</td>
<td>0.594</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sample Group III</td>
<td>0.75</td>
<td>-8.79</td>
<td>0.577</td>
<td></td>
<td></td>
<td>64</td>
<td>± 4</td>
</tr>
<tr>
<td></td>
<td>0.81</td>
<td>-8.78</td>
<td>0.596</td>
<td>0.579</td>
<td>0.016</td>
<td>0.6</td>
<td>± 1.2</td>
</tr>
<tr>
<td></td>
<td>0.79</td>
<td>-8.71</td>
<td>0.565</td>
<td></td>
<td></td>
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<td></td>
</tr>
</tbody>
</table>

For each sample the three Δ47 replicate measurements are averaged and show one standard deviation close to that found for homogeneous carbonate standards (±0.014‰, 1 SD). Uncertainties on temperature estimates are calculated from this ±0.014‰ external reproducibility or the SD obtained here if higher. Uncertainties on $\delta^{18}$Owater values reflect the variability in $\delta^{18}$O_carb and Δ47 values.

Figure 7: Relative timing of the diagenetic and fluid flow events affecting Chalk deposits in Flamborough Head (UK).
principal fluid flow event that led to the brecciation and extensive calcite mineralization (Figure 7).

**Burial Diagenesis and Matrix Cementation.** Within chalk deposits, burial diagenesis resulted in pressure solution processes taking place along stylolitic planes. Safaricz and Davison [66] estimated that 40% of the initial volume of chalk was lost due to chemical compaction. They estimated that dissolution of 30 mm thick of chalk results in a 1 mm thick of insoluble residue seam. A large number of marlseams and stylolites imply that 100s m of chalk was dissolved from the whole Coniacian-Campanian chalk. Chalk insoluble residue of Coniacian-Campanian chalk is lower than 5% so a high amount of dissolution is required to accumulate few millimeters of insoluble residue. The dissolution resulted in reprecipitation within the micritic matrix, as interparticle cement, thus affecting matrix porosity and permeability, with respective values as low as 8% and 0.04 mD [67]. Compaction stylolite planes, resulting from pressure solution, do not display any evidences of fluid circulation such as calcite cementation. In addition, δ18O of the sea from the ice-free Late Cretaceous sea is estimated at −1.2‰ [68]. The calculated δ18Owater of the fluids from which calcite veins precipitated range from 0 to 1‰. This provides an additional argument to support the fact that chalk is probably not the aquifer providing the mineralizing fluid, backing up the solid radiogenic Sr-isotope signal. At such low temperature, however, the isotope fractionation is probably kinetic and equilibrium may not have been reached, thus bringing an uncertainty to the calculated data.

**Late Cretaceous Inversion: Fluid Flow and Hydraulic Fracturing.** A phase of basin inversion occurred at the end of the Late Cretaceous-Early Cenozoic [24]. It affected many basins in NW Europe, including the Sole Pit Basin and the Cleveland Basin (Figure 1). Postburial structural inversion, particularly in shallow burial basins and at basin margin, allowed the reintroduction of meteoric water into the Sherwood Sandstone [52]. Bedding-parallel hydraulic fractures were reported in the Lower Jurassic shales of the Cleveland Basin [23] and Mesozoic mudstones and shales from the Wessex Basin [69]. In both cases, the timing of their formation was related to the Late Cretaceous to Cenozoic tectonic inversion. In Selwicks Bay, the tectonic inversion led to folding and tilting of the stratigraphy [13]. Some normal faults were reactivated during the inversion. Along the south coast of Flamborough (e.g., South Landing) only a preferential E-W orientation of the polygonal fault system was reactivated during the NNW-SSE compression and transpression events (described by Starmer et al., 2013). Many of the faults oriented perpendicular to the compression were reactivated as vertical stylolites. In Selwicks Bay, crosscutting relationships demonstrated that the tilting essentially occurred between Group I and Group II veins. Successive vein generations were identified in the field and geochemical analyses revealed that they originated from the same kind of fluid. The consecutive fluid flow events could be related to pulsating phases of inversion, which started during the Campanian and ended with the Laramide pulse of intraplate compression during Paleocene-Eocene [70].

Fluid flow under high pressure led to the formation of cemented breccia resulting from hydraulic fracturing [17]. Part of the overpressure may relate to the tectonic stress regime at the time, involving both the Alpine Orogeny and the Atlantic opening [23]. Nevertheless, previous studies [23, 69] suggest that tectonic forces alone do not allow reaching pore pressures high enough to initiate hydraulic fracturing, and the role of source rock maturation and oil generation was demonstrated. Our petrographical study...
including both fluid inclusion analyses and SEM observations of the calcite cements, however, did not display any evidence of hydrocarbon generation.

The comparison of the trapping temperatures inferred from microthermometry and clumped isotope thermometry gives coherent results. This suggests that cementation occurred at about 1 km paleoburial depth, as petrographically attested by the crosscutting relationship with compaction stylolites observed in Group I and III veins. The preferential circulation of fluids along reactivated faults was evidenced in chalk. A preexisting fault was reactivated, resulting in vein cementation in the damage zone of the basement fault and the polygonal fault system connected to it. Knowing that paleosealed breccia may cement fault zones in the subsurface is important to predict fault permeability and to identify potential drilling hazards in tight chalk.

6. Conclusion

A petrographical, geochemical, and microthermometric analysis of the calcite veins in the chalk of Flamborough Head (UK) was performed. The geochemical nature of the fluid and the relative timing of fault-controlled fluid flow have been constrained. These new data provide keys to understand the behavior of overpressured fluid flow in fractured tight chalks. The main conclusions are as follows:

(i) Mineralizing fluids are dominantly meteoric in origin. Fluid flow was probably triggered by Late Cretaceous-Cenozoic tectonic inversion and fluids circulated upward along the damage zone of the major fault.

(ii) Strontium isotopes demonstrate that mineralizing fluids probably originated from a sandstone aquifer (possibly the Triassic Sherwood Sandstone based on the local stratigraphy).

(iii) Fluid salinity locally increased through halite dissolution or fluid mixing with connate waters due to the interaction with lithologies overlying the aquifer.

(iv) In tight chalk, reactivated polygonal fault system was used as preferential fluid flow pathway at different stages of the geological history.

(v) Estimated crystallization temperatures obtained from both microthermometry and clumped isotopes range between 50 and 65°C.

Disclosure

An earlier version of this work was presented at “International Meeting of Sedimentology,” 2017.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

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References


