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Diversity and origin of quartz cements in continental carbonates: example from the 
Lower Cretaceous rift deposits of the South Atlantic margin

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Abstract

Silica precipitation in continental carbonates is a common process occurring during 
sedimentation and diagenesis. The Lower Cretaceous rift deposits of the South Atlantic 
equatorial margin, which are intensively explored by petroleum companies, provide good 
examples of such silicifications in carbonates, exhibiting a wide diversity of petrographic 
habit of early to late quartz cements. In order to understand the palaeoenvironmental and 
diagenetic conditions leading to this diversity, we integrated detailed petrography of 
diagenetic sequences and quartz habit with δ18Oquartz measurements (by SIMS) of individual 
cements observed in samples from the offshore and onshore basins of the West African 
margin. The petrographic description highlights the omnipresence of early fibrous 
microquartz cements exhibiting either length-fast or length-slow habit, in addition to 
laminated microquartz and micro- or mega-quartz forms. Amongst the isotopic analysis, the 
δ18Oquartz data show that length-slow cements are generally strongly enriched in 18O (δ18Oquartz 
ranging from 31 to 37‰ SMOW), whereas length-fast forms show less elevated values
(<32‰ SMOW). The highest $\delta^{18}O_{\text{quartz}}$ values for fibrous microquartz are interpreted to reflect precipitation from evaporated meteoric fluids at temperatures >25°C and <100°C. The alkalinity required to favor the precipitation of length-slow fibrous microquartz cements is probably related to fluid/rock interactions with underlying mantle-related or basic volcanic rocks. Such interactions would be in agreement with the recent geodynamic models of the South Atlantic passive margin. The length-fast fibrous microquartz associated with $\delta^{18}O_{\text{quartz}}$ values ranging from 27 to 32‰ SMOW, probably reflect precipitation from moderately to non evaporated, fairly neutral to acid, fluids. The partial dissolution of carbonate cements prior to quartz cementations represents the signature of those acidic conditions. We therefore suggest that acidic pH was obtained through fluid/rock interactions with the intermediate to acid volcanic rocks encountered along the palaeohydrological pathway. Other quartz phases, such as the megaquartz cement, exhibit highly variable $\delta^{18}O_{\text{quartz}}$ values ranging from 20 to 40 ‰. This variation may reflect significant variation in temperature conditions (between 100 and 200°C) or changes in fluid $\delta^{18}O$ at the small scale. For these populations of non-fibrous quartz cement, the very high $\delta^{18}O_{\text{quartz}}$ values may reflect a contribution of fluids that have either suffered strong evaporation or strong water/rock interaction.

Keywords: Quartz; Continental carbonates; Diagenesis; Fluid/Rock interaction; Oxygen isotope; Presalt

1. Introduction

Silicification of continental carbonate rocks is common during sedimentation and early to late diagenetic processes (e.g. Spötl & Wright, 1992; Bustillo, 2010; Alonso-Zarza et al., 2011). The relative commonness of these silicifications likely results from several factors. The first one may be related to the fact that continental carbonates are usually less widespread than marine deposits. Consequently, these deposits may remain connected with their silica-rich
substratum and surrounding catchment, from which alteration potentially provided dissolved silica. This abundance can also be due to potentially high silica contents in lacustrine, palustrine, or sinter settings, in relation to diatoms or as amorphous gels (natural gels in recent settings have only rarely been reported; Ernst & Calvert, 1969; Knauth, 1994; Lynne et al., 2005). Continental carbonate silicifications such as sinters may also occur in volcanic settings, associated locally with hydrothermal activity. Volcanic rocks are likely to release silica during their alteration (Hesse, 1989; Teboul et al., 2017). Finally, silicifications are extensively found in evaporitic settings, such as the Magadi Lake (Kenya; Eugster, 1967, 1969; Eugster & Jones, 1968).

In comparison to marine carbonates, quartz in continental carbonates are petrographically highly diversified. The most representative silica succession associated with cementation and/or replacement of carbonate or silica precursors, is typified by: (1) amorphous silica (opal-A), (2) opal-CT, (3) fibrous microquartz, and (4) megaquartz (Folk & Pittman, 1971; Marin, 2009; Warren, 2016). These sequences are seldom observed in marine carbonates, where epigenesis of micrite/allochems and replacement of evaporite nodules by microquartz constitute the majority of the indexed cherts, with other quartz morphologies occurring preferentially in large voids and fractures (Meyers, 1977; Hesse, 1989). Fibrous microquartz cements precipitating in primary and secondary pores of continental carbonates are very common. However, the context of their precipitation is less studied than megaquartz cements (Marin-Carbonne et al., 2011). Fibrous microquartz cements can form through ageing of previously deposited opaline minerals, or direct precipitation from silica-saturated fluids (Bustillo, 2010). They can be found under length-fast or length-slow forms (Folk & Pittman, 1971; Bustillo, 2010; Warren, 2016). Length-slow fibrous microquartz (or lutecite and quartzine) shows a crystallographic c-axis parallel or inclined by circa 30° to the fibers. Length-fast fibrous microquartz (or chalcedony), shows its crystallographic c-axis
perpendicular to the fibers. This interesting duality has been addressed by Folk and Pittman (1971), who used it to identify environments of precipitation. They suggested that length-fast cements are consistent with neutral to acid pH environments, whereas length-slow forms characterize alkaline environments. So far, the use and interpretation of the length-slow vs. length-fast criteria is mainly based on their petrographic features and their frequent association with evaporitic minerals. A recent approach from Bustillo et al. (2017) showed the predominance of length-slow forms in carbonate-evaporite precipitating lakes, and the predominance of length-fast forms in carbonate precipitating lakes. No studies have yet documented whether the isotopic composition of the different fibrous microquartz ($\delta^{18}O_{\text{quartz}}$) could potentially help to characterize their conditions of precipitation.

The present study focuses on the Lower Cretaceous deposits of the South Atlantic petroleum provinces (Carminatti et al., 2008; Beglinger et al., 2012; Saller et al., 2016). These deposits are overlain by a thick and extensive upper Aptian salt formation. Consequently, the deposits preceding salt deposition are commonly referred-to as “presalt deposits”. The most commonly prospected reservoir formations include the coquina succession (skeletal carbonate; Thompson et al., 2015), microbialites, spherulites, and calcitic shrubs (radial calcite), and possible hydrothermal carbonates (Terra et al., 2010; Saller et al., 2016; Szatmari & Milani, 2016). These deposits are interpreted as mostly derived from freshwater to hypersaline lakes (Saller et al., 2016). Little information has been published on the diagenesis of the presalt carbonates, and the available literature mainly focused on carbonate cementation and neomorphism (Frixa et al., 2015; Wright & Barnett, 2015; Gindre-Chanu et al., 2015, 2016; Saller et al., 2016). Despite the reported occurrence of large-scale quartz replacements and cementations (Wright & Barnett, 2015; Saller et al., 2016; Girard & San Miguel, 2017; Goldberg et al., 2017; Poros et al., 2017) the parameters triggering quartz cementation are still debated. Specific geochemical and crystallographic characterization has
been poorly developed on fibrous microquartz cements, with the exception of few $\delta^{18}$O$_{\text{quartz}}$ values from a single presalt site recently published by Saller et al. (2016).

The aim of this paper is to characterize and interpret various silica cements from several sites of the Angola and Congo margins through a combined petrographical and geochemical approach. The objectives of our study were therefore to characterize the petrographic habit of quartz cements, to constrain their relative timing of precipitation, and to characterize their $\delta^{18}$O$_{\text{quartz}}$ signature and the possible correlations with the petrographic characteristics. Finally, our aim was also to understand the relationship between the lithological heritage (i.e. substratum types in the palaeohydrological basins) and quartz precipitation, in light of recent works focused on the geological setting (Comin-Chiaramonti et al., 2011), the geodynamic evolution (Unternehr et al., 2010; Péron-Pinvidic et al., 2015), and the alteration of presalt volcanic substratum (Teboul et al., 2017).

2. Geological setting and studied material

2.1. Structural context

The onshore Namibe, Kwanza, and Lower basins were formed during the Neocomian rifting related to the breakup of Pangea (Pletsch et al., 2001; Szatmari & Milani, 2016). This asynchronous opening in time and space led to the initiation of four segments (equatorial, central, southern, and “Falkland”) bounded by major oceanic fracture zones (Torsvik et al., 2009; Moulin et al., 2010; Beglinger et al., 2012; von Nicolai et al., 2013). The Namibe, Kwanza, and Lower Congo basin are located in the Central Segment, aligned along a north-east to south-west transect, on the Angolan and Congolese coasts (Aslanian et al., 2009, Harris, 2000; Fig. 1A). The Namibe basin is bound to the south by the Walvis Ridge (Fig.
1A), which is a volcanic edifice developed above the Tristan da Cunha plume (Moore et al., 1984; O’Connor & Duncan, 1990). Rifting in the Kwanza Basin started at ≈144-140 Ma, with the onset of seafloor spreading at ≈127-117 Ma (Brice et al., 1982; Teisserenc & Villemin, 1990; Guiraud & Maurin, 1992; Quirk et al., 2013). Following Nürnberg & Müller (1991), seafloor spreading was active south of the Walvis Ridge at ≈126 Ma. According to Dingle (1999), during the early Cretaceous, the Walvis Ridge constituted a regional barrier to the marine conditions prevailing in the south.

According to Beglinger et al. (2012), the pre-rift basins are considered as intracratonic sag basins, whereas the syn-rift basins are associated with interior fracture basins, controlled by faulting and subsidence. The rifting led to the development of an extensive system composed of half-graben and horst. From the Berriasian to the Barremian, the fault-controlled subsidence led to the development of several fluvial to lacustrine basins filled from fault shoulders erosion products (Bate, 1999). In the early Aptian, the basins underwent thermal subsidence (sag), leading to the widening of the lacustrine systems and to the erosion of the uplifted fault blocks (Karner & Driscoll, 1999).

Recent studies aimed to establish new and more reliable models of the geodynamic evolution of intracratonic rifting and seafloor spreading (Moulin et al., 2010; Unternehr et al., 2010; Chaboureau et al., 2013; Péron-Pinvidic et al., 2015). They demonstrated a strong dissymmetry along the Angola-Namibe rifted margin, and pointed out that crustal thinning is probably favoured by the expansion of large detachment faults generating upper and lower plate domains (Unternehr et al., 2010; Péron-Pinvidic et al., 2015). Large basins with little subsidence would occur in the lower plate domain, controlled by ductile deformation. Narrow subsiding basins with abrupt margins would occur in the upper plate domain, associated with brittle deformation. These authors have localized the volcanic activity in the upper plate
domain where the thermal gradient is strong, thus inducing a strong vertical uplift associated with strong erosion and hydrothermal processes. Following Péron-Pinvidic et al. (2015), the south of the Lower Congo Basin and the Namibe Basin are associated with the upper plate domain. The Kwanza Basin is associated with a lower plate domain, but the studied wells are located close to transfer zones dividing the extending crust into segments. These zones can be either sharp (major normal crustal-scale fault) or diffuse (vague crustal and stratigraphic geometries; Péron-Pinvidic et al., 2015), and infer possible complex connections between lower and upper plate domains as well as crustal fluid circulations. According to Denis and Kluska (2017), the study area is located near the Benguela Transfer zone, where the extension is highly oblique and dissects the margin in narrow stripes, offsetting structural domains apart from transverse structures, superimposing two structural fabrics.

### 2.2. Palaeoclimate, stratigraphy and volcanism

According to Föllmi (2012), the most important factor controlling the Early Cretaceous climates is the progressive break up of Pangaea. He established the relationships between the climate type/intensity/evolution, the tectonic and volcanic activity, the continental vegetation cover and the variations in atmospheric CO$_2$ partial pressure. These proxies interacted with greenhouse conditions and generated an oscillation between arid and humid conditions. Thompson et al. (2015) emphasized the climatic disparities along the proto-South Atlantic rift.

Presalt basins were associated with continental-lacustrine settings during syn-rift phase (Guiraud et al., 2010). Pre-rift and syn-rift sediments are mainly composed of clastic and carbonate deposits reflecting continental alluvial, fluvial, and lacustrine environments (Beglinger et al., 2012; Chaboureau et al., 2013; Fig. 1B). The early Aptian thermal subsidence favoured the development of restricted brackish to hypersaline areas as indicated
by the deposition of coquinas, travertine and microbial carbonates on isolated platforms (Gomes et al., 2009; Thompson et al., 2015). The presalt carbonate and clastic deposits are overlain by widespread and massive Aptian evaporites (Karner et al., 2003; Karner & Gambôa, 2007), known as the Loeme Formation in the Kwanza and Congo basins (Chaboureau et al., 2013), and as the Bambata Formation in the Namibe Basin (Gindre-Chanu et al., 2015, 2016). Chaboureau et al. (2012) concluded that those evaporitic systems were partly fed by hydrothermal recharges. A complete review of the paleogeographic and stratigraphic evolution of the presalt is available in Chaboureau et al. (2013).

A widespread magmatic activity occurred coevally to presalt deposits (Pereira, 1969; 1971; Marzoli et al., 1999; Comin-Chiaramonti et al., 2011, Teboul et al., 2017). The magmatic budget changed from south to north, with magma-rich provinces in the south and magma-poor provinces in the north (Péron-Pividic et al., 2015). A synthesis of the magmatic setting of the Kwanza Basin is available in Teboul et al. (2017). Comin-Chiaramonti et al. (2011) have shown that the Namibe Basin volcanic rocks follow similar trends than those from the Kwanza Basin. Gindre-Chanu et al. (2016) linked the volcanic rocks occurring in the Namibe Basin to the Bero Volcanics Formation, which overlie the basement. These mainly consist of tholeiitic to transitional basalts and rhyodacites interlayered with volcaniclastic and aeolian to fluvial sandstones. Likewise, the south part of the Kwanza basin registered non-negligible syn-rift and early post-rift volcanism (Aptian in age) from tholeiitic to transitional (Denis & Kluska, 2017) encountered in a number of Angola offshore wells. Little information is found in the literature on the occurrence of Cretaceous magmatism in the Congo Basin. Jackson et al. (2000) argue that unpublished industry seismic data indicate that volcanic rocks probably occur in this basin. Giresse (2005) reported the occurrence of kimberlitic intrusions during Cretaceous time.

2.3 Studied material
Confidentiality restrictions preclude the disclosure of the exact locations, names, and depths of the wells.

In the Lower Congo Basin, the studied samples come from two cored offshore wells (Toca-1 and Toca-2) that belong to the synrift carbonate reservoirs of the Toca Formation. This formation is intercalated between and within two source rocks, the Pointe Noire and Pointe Indienne shales (Chaboureau et al., 2013). The Toca Formation (short for “top carbonate”) is dominated by skeletal carbonates with a main facies characterized as “coquina” facies (Harris, 2000; Thompson et al., 2015), which are presently studied from the Toca-1 and Toca-2 cored wells. This facies is characterized almost entirely by coarse (<2 cm) packstone to grainstone composed of unionid bivalves and gastropod shells, with occasional ostracod and fish remains. As stated by Thompson et al. (2015), coquinas were formed in freshwater to brackish lakes, at shallow depths, on horst blocks, hangingwall ramps and footwall crests of tilted fault blocks, accommodation zones, and anticlinal rollover structures.

The studied material from the Kwanza Basin (offshore Angola) has been collected on sidewall cores of the Shrubby-1 well, and from a core of the Berilo well. In Shrubby-1, part of the well has been assigned to the Cuvo Formation, composed of syn-rift fining upward litharenite/conglomerates (Bate et al., 2001). These clastic deposits are mainly overlain by dolomitic and silica-cemented carbonates and evaporites, considered as an equivalent of the Chela Formation in this study. This formation has been studied in the upper part of the well Shrubby-1 and in Berilo. The Cuvo and Chela Formations are associated with the thermal subsidence preceding salt deposition, which developed on a regional stratigraphic intra-Aptian discontinuity (San Miguel et al., 2017). Carminatti et al. (2008) also showed that these carbonates accumulated on structural highs, which are currently located in deep-water settings offshore Angola (>1000 m below sea-level).
Two outcrops from Mariquita and Mina do Gesso (Onshore Namibe Basin, Angola) have been studied. These outcrops are respectively located 2.5 km east and 9 km north-east of the Mariquita village. Presalt deposits of the Namibe Basin are poorly described in the literature (Coterill et al., 2002; Chaboureau et al., 2013; Gindre-Chanu et al., 2015, 2016; Schröder et al., 2015). According to Gindre-Chanu et al. (2015, 2016), lacustrine marlstones, limestones, and microbial deposits belong to the Cangulo Formation. These microbial deposits are associated with dolomitic and siliceous bodies attributed to syn-sedimentary hydrothermal activity. The overlying formation, called by Gindre-Chanu et al. (2015, 2016), the “Tumbalunda Formation” consist of fluvial to alluvial fan gravelly facies, conglomeratic mouth deltaic bars, metre-thick siltstones, organic-rich mudstones, siliciclastics, evaporites, and carbonate beds formed either in lacustrine or hydrothermal setting.

3. Methods

3.1. Paragenetic sequences

Conventional optical microscopy, under plane polarized light (PPL) and cross-polarized light (XPL), was used to examine 300 polished thin sections in order to characterize microfacies and diagenetic phases such as cements, dissolution phases, fractures, and mineralogical replacements. For calcite and dolomite identification, thin sections were partially stained with alizarin red-s and potassium ferricyanide after etching in dilute hydrochloric acid (Dickson, 1966). A first order (full wave) retardation plate has been used in order to characterize the sign of elongation (length-fast or length-slow) of fibrous microquartz cements. Additional methods were applied to selected samples. These methods include the use of cathodoluminescence (referred as “CL”) on a Luminoscope ELM-1 coupled with a stereomicroscope Leica MZ12 and ZEISS MRc5 camera (Burgundy University) and a CCI 8200 Mk4 coupled with a Nikon Eclipse 80i microscope and a Nikon DS-Fi1c camera.
UV epifluorescence microscopy was carried out on a AZ100 Nikon macroscope under a 360 nm excitation LED light. Fibrous microquartz microtextures were investigated on six rock chips using Scanning Electron Microscopy (XL30 SEM™; FEI/Philips; Université Aix-Marseille). A total of twenty-two thin sections were analysed on a Scanning Electron Microscope equipped with a Brucker Quantax EDS detector (SEM-Quantax; TOTAL CSTJF laboratory), allowing to investigate the elemental composition of specific minerals. Microthermometric measurements of fluid inclusions were carried out, when possible, using a THMS G600 Linkam semi-automatic gas-flow freezing-heating stage (Université Bourgogne-Franche Comté).

3.2. Oxygen isotopes

In situ oxygen isotope microanalysis of quartz cements was performed by secondary ion mass spectrometry (SIMS) at the CRPG in Nancy (France) using a Cameca 1280HR. Analyses of different samples were performed over six analytical sessions in two periods of three successive days (July 20\textsuperscript{th}-21\textsuperscript{st}-22\textsuperscript{nd}, 2015 and May 30\textsuperscript{th}, 31\textsuperscript{st}, June 1\textsuperscript{st}, 2016). Analyzed thin sections were finely polished (with 0.25 μm diamond), then checked for flatness and photographed using a reflected light microscope. This was done to facilitate the precise location of areas of interest (the SIMS 1280 is devoid of transmitted light visualization system) and especially to avoid areas with reliefs that are known to substantially modify measurements (Eiler et al. 1997; Valley and Kita 2009). A total of 19 thin sections from the different studied locations have been analyzed. A Cs+ beam with a 10 kV accelerating potential is used to sputters sample atoms, excavating a shallow pit in the surface (~5 μm in diameter). The $^{18}$O and $^{16}$O isotopes were measured on two off-axis Faraday cups, and $^{17}$O isotopes on the axial electron multiplier in multicollection mode. Measurements were performed with typical counts of $\sim 3 \times 10^9$ counts per second for $^{16}$O. Acquisition sequences
consisted of 60 s of pre-sputtering and 30 cycles resulting in an internal error better than 0.1‰ ($2\sigma_n$, $n =$ number of cycles). The external reproducibility was estimated through multiple analyses of two standards, with systematic measurements at the beginning and at the end of each session (days). The two laboratory standards (QZ-BRA and NL 615; Fouillac and Girard, 1996; Marin et al., 2010) are homogenous megaquartz (Girard et al., 2001) as no opal or fibrous microquartz standards are available. They are included in epoxy, finely polished, and mounted in a single cylindrical mount. For each session, the drift between first and last standard measurements was calculated, leading to an external reproducibility of $\delta^{18}$O from $\pm 0.1$ to $\pm 0.6$ (1$\sigma$). QZ-BRA has an accepted value of 9.6±0.2‰ and NBL 615 has an accepted value of 18.4±0.2‰. Eiler et al. (1997) suggested that the analysis of a standard with a chemical and structural composition similar to the studied samples is needed to apply proper correction of the matrix effect (instrumental bias). Areas with epoxy-impregnated microporosity were not measured, or were not integrated in the results, in order to avoid the possible influence on this epoxy on the oxygen isotopic signal. All $\delta^{18}$O values are expressed in permill relative to the Standard Mean Ocean Water (SMOW).

4. Results

4.1. Petrography and cement stratigraphy

The diagenetic sequences depicted in figure 2 are based on textural relationships observed in thin sections. In agreement with the scope of this study, the following paragraphs focus on the main diagenetic stages and the description of silica cements.

4.1.1. Onshore Angola

In Mariquita, the primary facies is laminated, consisting of a succession of (i) inclusion-rich radiaxial calcite layers, (ii) peloidal wackestone to grainstone and (iii) thin micritic laminae.
All these layers are intensively, but not systematically, dolomitized. When not dolomitized, the micrite is Fe-poor and non-luminescent. When dolomitized, the matrix is cryptocrystalline and has a dull to bright red luminescence. Radial layers are now composed of Fe-poor and inclusion-rich dolomite, but various characteristics suggest a primary calcitic nature, as highlighted by the striking similarity with the calcitic radial layers and spherulites observed in Mina do Gesso outcrops. This dolomite shows a sweeping extinction with a dull-red luminescence. Botryoids of dolomitized calcite are also observed in early fractures and form the first cements. The latter are overlain by euhedral dolomite rhombs with a dull red to brown CL and a low Fe content. Silica precipitated after a phase of fracturing and strong dissolution of calcite and dolomite (Fig. 2). In few samples, the first silica cements are isopachous or botryoidal length-fast fibrous microquartz. In most samples, silica cements are only represented by megaquartz cements (up to 1mm long) that precipitated either in primary pores, in vuggy pores, in fractures, or as a replacement of primary radial calcite layers (Fig. 3A, B, C, D and E). Occasional later cements are either micritic pendant cements or Fe-poor blocky calcite with a dark red luminescence. The latter are probably of late vadose origin.

The facies studied in Mina do Gesso area are essentially composed of microbial mats displaying micritic clotted textures, with occurrences of inclusion-rich radial calcite forming discontinuous layers and spherulites (Fig. 3F). Cements developed before compaction, mainly in primary fenestral porosity. The first cement is euhedral Fe-rich dolomite rhombs followed by two zones of blocky calcite displaying an orange luminescence (Fig. 2). The first one is Fe-rich and the second one is Fe-poor. Some dolomite rhombs underwent dedolomitization. Silica cements developed latter, after a phase of partial dissolution of calcite and dolomite (Fig. 2, dissolution “d1”). Most are length-fast fibrous microquartz that seals the remaining porosity (Fig. 3G). Anhedral megaquartz cements also
replace radiaxial fibrous calcite (Fig. 3F). The timing of these replacements is poorly constrained. Iron oxides, probably related to telogenesis, postdate all cements.

For both areas, an accurate quantification of quartz precipitation is particularly difficult as a sampling bias is likely influencing our observations. At outcrop scale, Mina do Gesso is poorly silicified in contrast with the extensive quartz silicification of the Mariquita section.

4.1.2. Offshore Kwanza

Two diagenetic sequences are observed in Shrubby-1 sidewall core samples and are associated with the Chela Formation and the underlying Cuvo Formation:

- The Chela Formation is composed of fibrous radiaxial calcite (*i.e.* spherulites) and shrubs-like components (Fig. 4A), which occasionally are dolomitized. Additionally, small euhedral dolomite rhombs with a dull red luminescence occur in the intergranular porosity. The shrubs, spherulites, and dolomite rhombs are enclosed in several silica cements (Fig. 2; 4A, C). From overall point of view, extensive silica precipitations occur along 93 m of the cored section, with a maximum of 95% of silicification at some stratigraphic level. The first silica cement occurs as length-slow radiaxial fibrous microquartz forming isopachous fringes and botryoids up to 500 µm thick. This cement is named porous fibrous microquartz (pfQ1) due to its frequently porous microstructure detectable both under PLL and SEM (Fig. 4 A, B, D, G, I). In PPL, the porous habit results in a blue colour due to the presence of blue epoxy resin between fibers (Fig. 4A & D). In other cases, this microporosity has been partially filled by oil, now transformed into bitumen, displaying brown patches at the periphery of pfQ1 cement (Fig. 4C). Under SEM, the fibers of pfQ1 are composed of aligned very small silica crystals exhibiting an anhedral shape and a diameter < 0.25 µm. These small crystals alternate with coarser subhedral quartz crystals (up to 3 µm long)
having their c-axis parallel to the fibers of the cement (Fig. 4G). In most thin sections, the pFQ1 cement is followed by 10 to 50 \( \mu \text{m} \)-thick isopachous cement characterized by alternations of equigranular microquartz and fibrous microquartz laminae (LQ1; Fig. 4). The latter is followed by a second generation of radiaxial fibrous microquartz cement with a petal like texture (referred as “petaloid”) and a brown colour under PPL (pFQ2; Fig. 4D). pFQ2 was extremely microporous initially, but the microporosity between fibers is now filled with bitumen, as indicated by its brown colour (Fig. 4D). This suggests a probable syn-pFQ2 oil charge (see discussion). A new generation of 10 to 20 \( \mu \text{m} \)-thick isopachous laminated microquartz cement (LQ2; Fig. 4D) commonly overlaps the LQ2 petaloids. Finally, equigranular megaquartz cement (MQ), composed of euhedral colourless inclusion-poor crystals with a size up to 400 \( \mu \text{m} \), fills up the remaining porosity (Fig. 4A, B, H). These successive quartz generations are considered as precompaction cements based on the evident lack of compaction features between the carbonate grains. Several fracturing stages have been observed. The first generation of fractures occurs after LQ1 and before pFQ2 precipitation. A second generation of fractures occurs before megaquartz cementation. Several fractures are also associated with microquartz cementation, but their timing is hardly constrained due to the lack of crosscutting relationships. Two successive dolomite cement phases seal a new generation of fractures associated with local calcite dissolutions: the first one shows planar euhedral crystals with dull red luminescence, whereas the second one corresponds to saddle dolomite crystals with red luminescence. These cements occur either in the remaining intergranular porosity, in fractures, or in dissolution voids located in the former shrubs. Finally, the remaining porosity is plugged by blocky calcite with a yellow luminescence, and by
bitumen. This last occurrence of bitumen is probably related to a late oil charge, based on the oil impregnations observed in the porous fibrous microquartz.

- In the Shrubby-1 well, the underlying Cuvo Formation is composed of siliciclastic deposits, basaltic beds and gabbroic bodies. The diagenetic features described below are mainly located in fractures. Firstly, the fractures are partially filled by dolomite cement, with euhedral crystals larger than 250 µm and a dull red luminescence (Fig. 2). Dolomite crystals sometimes show undulose extinction under XPL. They are enclosed in several successive generations of quartz cements starting by porous fibrous microquartz cements (pFQ) forming thick isopachous fringes or botryoids. These cements are highlighted by the blue epoxy colour under PPL. This microporosity is sometimes partially filled by oil, now transformed into bitumen. Under XPL, this cement is recognizable by its distinctive radiaxial extinction. pFQ is followed by two generations of megaquartz cements (MQ1 and MQ2) alternating with two generations of 10 to 500 µm-thick isopachous laminated microquartz cements (LQ1 and LQ2; Fig. 5A, B). As in the overlying Chela Formation, the laminated microquartz fringes are defined as alternating equigranular microquartz laminae and fibrous microquartz laminae. MQ cements shows euhedral palisadic to equigranular colourless quartz crystals larger than 400 µm in size (Fig. 5A, B). The second megaquartz cement (MQ2) is enclosed in blocky calcite cements displaying a yellow luminescence. The remaining fissural porosity contains occasional traces of bitumen.

In the Berilo well, quartz cements are observed in an intensively dolomitized facies showing ghosts of microbial laminae with fenestrae, thrombolitic micritic textures and peloidal cloths. Small subhedral dolomite crystals (< 5 µm) with an orange-red luminescence replace the initial deposits (Fig. 2). Associated euhedral dolomite crystals, < 150 µm in size, border the fenestral pores. Under PPL, this dolomite shows a cloudy
inner zone and a clean, limpid outer zone. One two-phase liquid-rich secondary fluid inclusion occurring in the outer zone provided a homogenization temperature of 109 °C. These cements are enclosed in authigenic barite, occurring as individual crystals, or combined in radiating sheaves. Many two-phase liquid-rich secondary fluid inclusions (water/gas) occurring in the barite were analyzed, yielding homogenization temperatures ranging from 50 to 120 °C. The fenestral porosity is finally totally plugged by two quartz cements i.e. botryoidal length-fast fibrous microquartz with a flamboyant texture (Fig. 5C) and a mosaic of euhedral to subhedral megaquartz crystals, ranging in size from 50 to 250 µm (Fig. 5C). Quartz cements represent around 3% of the studied facies of this well. Unfortunately, no fluid inclusions have been observed in this latter cement.

4.1.3. Lower Congo Basin

The coquina facies of the two wells studied in the Lower Congo Basin are characterized by first cements consisting of blocky calcite crystals (BC1) showing a scalenohedral habit and a succession of dull to yellow bands under CL. These calcite crystals are Fe-poor, Mg-poor and inclusion free. They developed both in interparticle and moldic pores due to an early partial dissolution of bivalve and gastropod aragonite shells. The precipitation of BC1 occurred in very shallow burial conditions, as indicated by the lack of any compaction feature. When not dissolved, the biogenic aragonite is replaced by a neogenetic inclusion-rich calcite (AC; Fig. 2) showing anhedral growth bands typical of the neogenetic replacement process. Many two-phase liquid-rich inclusions (water/gas and oil/gas) have been found and analysed in AC, indicating homogenization temperatures ranging from 70 to 180 °C and from 60 to 120°C in the Toca-1 and Toca-2 sections respectively. The secondary origin of these fluid inclusions is highlighted by their location along cleavage planes, as well as sealed microfractures. In Toca-1 samples, a second generation of blocky calcite zone (BC2; Fig. 2) took place before silica
precipitation, but after the formation of microfractures, indicating mechanical compaction. This orange to brown CL zoned cement grew syntaxially on BC1. It consists of a Fe-rich calcite as revealed by alizarine-ferricyanide staining and SEM-EDS observations. BC2 contains two phase, liquid-vapour, inclusions mainly located along cleavage planes. Microthermometric measurements in these secondary fluid inclusions provided homogenization temperatures ranging from 80 to 165 °C.

Silica cements in Toca-1 and Toca-2 precipitated after a weak dissolution of BC1 (Toca-2) and a strong dissolution of BC2 (Toca-1) cements, as revealed by dissolution vugs plugged by silica cements. Microfractures also affect shells and BC1+BC2 cements but not the silica cements. This suggests that the precipitation of silica cements occurred after mechanical compaction in the interparticle porosity (i.e. between the shells) and in the remaining moldic porosity (Fig 5D, E, F, G). In both wells, quartz precipitations are non uniform along the cored sections. In Toca-1, several silicification fronts are observed, with a maximum thickness of 10 cm, however in Toca-2, those fronts can reach 20 cm.

Successive fibrous microquartz and megaquartz cements are observed. In Toca-1 and Toca-2, the observations under PPL show a first generation of light to dark brown fibrous microquartz cement having an isopachous or a botryoidal outline. Under CL, these cements show some finely concentric laminae with alternating bluish to dark luminescence. A well-expressed sweeping extinction is detected under XPL. The use of the full wave retardation plate reveals that fibrous microquartz are length-slow in Toca-2, but length-fast in Toca-1. In the two cases, the fibrous microquartz are dense, with limited microporosity between fibers.

Megaquartz cementation mainly occurred in Toca-2, after a fracturing phase (Fig. 5F, G). These crystals display a sweeping extinction under XPL and alignments of solid inclusions
displaying chevron structures along growth bands. Neither biphasic liquid/gas inclusions nor oil-rich inclusions have been detected in these megaquartz crystals.

A single phase of blocky calcite postdating silica cementation is observed in Toca-1. This Fe-rich, blocky calcite has an orange colour under CL. In Toca-2, the diagenesis postdating silica cements is more complex and is composed of one generation of dolomite and two generations of blocky calcite. The dolomite rhombs are euhedral, Fe-poor, and exhibit three successive red to dark red zones under CL. In Toca-1 and Toca-2 the late cements consist of Fe-rich blocky calcite zones with a dark orange to brown luminescence. These late blocky calcites locally precipitated in fractures that crosscut silica cements. The remaining porosity is locally filled with bitumen.

4.2. Oxygen isotopes

The $\delta^{18}$O values measured by SIMS in the silica cements at the different studied sites are compiled in Figure 6 and are reported in a supplementary data file that contain both data and extra microphotographs (#supplementary data 1). A total of 203 values are measured, showing an extremely large range going from $\sim$20‰ to $\sim$40‰ and covering about 50% of the documented range of $\delta^{18}$O variation in rocks & minerals of sedimentary origin (i.e. 0-40‰). The microporous length-slow fibrous microquartz cements tend to show elevated $\delta^{18}$O values, ranging from 33 to 37‰ in Shrubby-1 (33 to 35‰ (N=4) in the Chela Formation and 33 to 37‰ (N=4) in the Cuvo Formation) and 31 to 32‰ (N=9) in Toca-2. The non-porous length-fast fibrous microquartz cements tend to show somewhat lower $\delta^{18}$O values, ranging from 28 to 31‰ (N=10) in Berilo, 29 to 32‰ (N=25) in Toca-1 and around 28‰ (N=4) in Mina do Gesso.
The laminated microquartz in Toca-2 shows very variable $\delta^{18}O$ values ranging from 27 to 33‰ (N=30). This is also true of the laminated microquartz in Shrubby-1, where $\delta^{18}O$ values range from 27 and 30‰ in the Chela (N=2) and 32 to 39‰ (N=8) in the Cuvo Formation.

The greatest overall variability in $\delta^{18}O$ values is observed in the megaquartz cements (Fig. 6), although values can be pretty consistent in a single site. $\delta^{18}O$ values range from 21 to 30‰ (N=38) in Mariquita, 30 to 34‰ (N=16) in Toca-2, 29 to 30‰ (N=5) in Berilo, and 20 to 40‰ in Shrubby-1 (33 to 40‰ (N=23) in the Chela Fm. and 20 to 38‰ (N=25) in the Cuvo Fm.) The highest $\delta^{18}O$ values of 39-40‰ measured in this study come from the megaquartz in the Chela Formation and are amongst the highest values ever reported for quartz. The lowest $\delta^{18}O$ values of 20-21‰ recorded in our dataset come from the latest growth stage of the megaquartz cement (Fig. 5B) in the Cuvo Formation. When possible, the small analytic spots (10–20 microns in diameter) were situated along transects in order to document any spatial-temporal variability in the $\delta^{18}O$ values of single cement (Fig. 3E, G; 5B, C, E). The only observed trend occurs in a Mariquita sample showing a $\delta^{18}O$ increase through time by a few ‰ (Fig. 3E).

5. Interpretation and discussion

5.1. Source of aqueous silica during presalt diagenesis

As stated by Hesse (1988, 1989), silicification is a common diagenetic process occurring in non-siliceous sediments. Several authors discussed the silica budget in oceanic (e.g. Heath, 1974; Riech & v. Rad, 1979; DeMaster, 1981; Hesse, 1988; Tréguet et al., 1995) and continental (e.g. Cornwell & Banahan, 1992; Arai & Fukushima, 2012; Pittari et al., 2016) settings. According to DeMaster (1981), some of the major sources of dissolved silica in marine waters are: (1) rivers, (2) active volcanism and associated hydrothermal emanations, (3) epigean weathering of volcanic rocks, and (4) dissolution of siliceous biogenic tests in
sediments. In lacustrine sediments, similar silica sources have been documented (Livingstone 1963; Renaut et al., 1998; Alexandre et al., 2004), but Bustillo & Alonso-Zarza (2007) also emphasized the possible contribution of the dissolution or alteration of clay minerals.

In Presalt rocks, dissolution of siliceous biogenic tests is not considered as a possible source of silica during diagenesis because marine organisms forming opal-A tests (radiolarians, siliceous sponges, silicoflagellates…) have never been observed in the continental to paralic sediments of the studied sites. Diatoms are also absent in the Presalt lacustrine sediments of the studied area.

A very likely source of dissolved silica in Presalt diagenetic fluids is the alteration of volcanic products (Tosca & Wright, 2015; Szatmari & Milani, 2016; Teboul et al., 2017). This alteration could have taken place either in catchment areas, in lakes, or in the substratum beneath the lakes through hydrothermal processes. A similar source of Si has been suggested for the continental rift basin of Limagne (Massif Central, France) where length-slow fibrous silica developed within lacustrine carbonate formations, in close association with brecciated volcanic material and bitumen (Devouard & Bornet, 2015). Cerling (1994) showed that the drainage of volcanics in surrounding catchments has a strong influence on the chemistry of the lakes of the African Rift Valley. It tends to highly increase alkalinity, producing dilute waters which chemical composition is dominated by calcium, magnesium, and silica. Furthermore, interaction of saline waters would be poor in alkaline earths (Cerling, 1994). In addition, the occurrence of syn- to post-sedimentary volcanism associated with hypogean to epigean springs feeding the lacustrine systems, may have induced high CO$_2$ input, high alkalinity, and high concentrations of dissolved silica, magnesium, and calcium (Wright, 2012; Teboul et al., 2017). Numerical modeling shows that the hydrothermal and meteoric alteration at variable temperature and CO$_2$ partial pressure of trachytes from a buried presalt
sequence favors high Si and Mg concentrations (Teboul et al., 2017). Presalt volcanism is characterized by bimodal, mafic and felsic, rocks including silica-saturated basalts and rhyolitic end members (Marzoli et al., 1999). Based on the abundance of calcite and Mg-silicates and on the absence of chlorides and sulphates, several authors highlighted that the weathering of basic volcanic formations are probably the main source of elements in presalt systems (Tosca & Wright, 2015; Szatmari & Milani, 2016). However, geochemical simulations designed to evaluate the exact contribution of the presalt basaltic volcanism to the diagenesis of Presalt deposits are yet to be conducted.

Several authors also described the influence of chemical weathering of volcanic debris and lavas in lacustrine systems as a source of solutes (e.g. Abdelouas, 1996; Garcia-Romero et al., 2005). The alteration products of glass-rich volcanic ash tend to produce Si-rich gels, Fe-rich oxy-hydroxides, clay minerals and zeolites (Abdelouas, 1996). Such alterations may have a direct impact on lake chemistry through the dissolution of volcanic glass, or an indirect impact through precipitation of “intermediary media” such as clays/zeolites. According to Millot (1970) in Tosca & Wright (2015), in close basin settings associated with alkaline fluids, Al- to Fe-rich clay minerals are located along the basin margins, where aluminosilicate detritus are abundant, while Mg- and Si-rich (and Al-poor) clay minerals are located in the center. Mg- and Si-rich clay minerals are common in presalt deposits on the Brazilian margin of the South Atlantic Ocean, in particular the phyllosilicate stevensite ($\text{Na}_{0.3}\text{Mg}_{2.7}\text{Si}_4\text{O}_{10}[\text{OH}]_2$; Thompson et al., 2015; Tosca & Wright, 2014; 2015; Wright & Barnett, 2015, Goldberg et al., 2017). Precipitation of Mg-silicates is favored at high pH ($\geq 8.7$) and Mg/Si ratio ($\geq 0.67$) in saline waters (Tosca and Wright, 2014; 2015; Zeyen et al., 2015). If altered, clay minerals could represent an important source of silica during diagenesis (MacKenzie et al., 1967; Thiry & Maréchal, 2001; Michalopoulos & Aller, 1995, 2004). Several authors emphasized that early to late diagenetic alteration of Mg-silicates
controlled, among other things, by pH or CO\textsubscript{2} partial pressure fluctuations, would result in the release of Mg, Si, and minor Na (e.g., Deocampo, 2005; Tosca & Wright, 2014; 2015). This alteration may have contributed to dolomitization and silicification processes in presalt deposits.

The actualistic comparison with published data reveals that many of the modern silica-rich waters are related to rift settings, and more particularly to lakes and springs of the East African Rift System (Eugster & Jones, 1968; Dunkley \textit{et al.}, 1993; Sturchio \textit{et al.}, 1993; Behr & Röhrich, 2000; Renaut \textit{et al.}, 1998, 1999, 2002; Dekov \textit{et al.}, 2014). Other case studies are usually associated with volcanic geological setting such as Specchio di Venere (Italy; Cangemi \textit{et al.}, 2010), Mono Lake (USA; Souza-Egipsy \textit{et al.}, 2005), Cachi Laguna and Pastos Grandes (Bolivia; Risacher, 1978; Abdelouas, 1996; Sylvestre \textit{et al.}, 2001), and Azores Islands (Brunet & Bustillo, 2014). Several authors attributed the occurrence of these silica-rich waters to the prevalence of closed-basin conditions and the primary inflow of alkaline sodium carbonate/bicarbonate springs associated with volcanic material (Kempe & Degens, 1985; Abdelouas \textit{et al.}, 1995; Abdelouas, 1996; Garcia-Ruiz, 2000). This agrees with the hypothesis advocated for presalt environments, stating that silica-rich fluids are derived from altered volcanic material. However, modern environments are often associated with diatom frustules. We consider that these frustules may act as silica-rich intermediary media, as the biomineralized silica is probably derived from the alteration of catchment areas.

Following the work of Ambwani \textit{et al.} (2003), the studied Early Cretaceous rocks are probably too old to use diatoms as intermediary silica source. In the modern deposits of Cachi Laguna (Bolivia), Abdelouas (1996) evidenced a close relationship between the occurrence of silhydrite (an hydrated form of silica: 3SiO\textsubscript{2}[H\textsubscript{2}O]), magadiite (hydrated sodium silicate) and stevensite-saponite, associated with the dissolution of nearby silicates,
rhyolitic volcanic glass, and diatom frustules. The latter author attributed the direct precipitation of silhydrite to low Na concentrations of hydrothermal waters, whereas magadiite tends to precipitate in Na-rich waters in the central part of the lake. Stevensite precipitation would be favored through the dissolution of the diatom frustules, in Mg-rich, alkaline setting, where silanol groups are available for clay precipitation. The Cachi Laguna area may therefore constitute a great location to establish the spatial and temporal relationships between volcanic material alteration, Mg-rich clay minerals and silica precipitation. In this case study, the potential of hydrated silica deposits to be remobilized or neomorphized during diagenesis should be investigated.

Finally, a recent study conducted by Pinto et al. (2016) evidenced that serpentinization of mantle rocks may also constitute a source of silica. These authors established mass balance calculations based on material from exhumed and serpentinized domains from Western Iberia and fossil Alpine Tethys margins. By extrapolation, they suggested that similar elemental transfers can occur in nascent oceanic basins, prior to seafloor spreading. Following the recent geodynamic models implying upper and lower plate domains, the exhumed mantle could have been coupled to presalt sediments through lithospheric-scale low angle detachment faults, enabling large-scale fluid circulation that may have influenced the presalt sedimentary-diagenetic evolution (Unternehr et al., 2010; Péron-Pinvidic et al., 2015).

In this study, the spatial distribution of quartz precipitation between the different sites is non uniform. The most extensive precipitations have been observed in the well Shrubby-1 and in the onshore outcrop of Mariquita. The vertical and lateral variation of silica content is probably related to differential amounts of silica supply during diagenesis of each studied area. Following the available knowledge of the presalt series, we consider that the most
probable sources of silica may be associated with the alteration of volcanic material or with fluids linked to mantle rock serpentinization.

5.2. Timing between carbonates and silica cementation

Some carbonate diagenesis predates silicification at all studied sites (Fig. 2). The earliest diagenetic processes include carbonate cementation and neomorphism, and are highlighted by: (i) a strong dolomitization and/or euhedral dolomite precipitation filling pores (Mariquita, Shrubby-1, Mina do Gesso, Berilo), and (ii) aragonite to calcite neomorphism of shells and blocky calcite cementations (Toca-1 and Toca-2). An early diagenesis controlled by carbonate precipitation is particularly consistent with presalt carbonate dominated sedimentary systems (Terra et al., 2010; Gindre-Chanu et al., 2016; Saller et al., 2016). Typical presalt carbonates are composed of shrubby microbial boundstone, and wackestone to packstone bearing calcitic spherulites in a dolomitic to argillaceous matrix (Terra et al., 2010; Saller et al., 2016; Goldberg et al., 2017). Such primary calcite (Mariquita and Mina do Gesso) and eventual primary dolomite (Berilo) structures are indeed observed in our studied material. This implies that a sedimentary-diagenetic continuum existed between the carbonate-rich lacustrine to paralic system and early diagenetic processes. The nature and succession of early diagenetic features support a continuum through precompaction cementation, the spherulites, and the cemented coquina. The fact that diagenetic silica comes later in the diagenetic succession, after a significant phase of calcite-dolomite diagenesis, constitutes an argument against the occurrence of sinter-like subaerial deposits at our studied sites, as silica would have represented the primary sedimentological and early diagenetic products. However, this observation does not preclude a relatively early formation of silica cements in the Mariquita outcrops and in the Chela Fm in the Shrubby-1 well, as in these two sites, silica cements precipitated before any significant mechanical compaction. This pre-
compaction diagenesis is exceptional in Mariquita, where the texture of primary carbonate deposits is particularly well preserved. Saller et al. (2016) also reported early pre-compaction silicification in the offshore Kwanza Basin, based on occurrence of intraclasts with truncated chalcedony cements interpreted as silica precipitation during spring mound-like deposition. This indicates that silica diagenesis started at the sediment/water interface or within shallow burial settings. In other studied sites, silicification may have occurred at a later time. In Mina do Gesso and Berilo, silica cement occurs in residual pore spaces that have survived a significant carbonate/sulfate cementation phase, suggesting silica precipitation took place during later diagenetic stages. Based on petrographic observations, quartz precipitation postdates the formation of microfractures which affect only shells and predates blocky calcite cementation in Toca-1 and Toca-2 studied sites. This suggests that quartz cementation occurred after the first steps of mechanical compaction. Finally, the silica cements of the Cuvo Fm. in the Shrubby-1 well occur mainly as fracture filling, cross-cutting moderately to strongly indurated sandstones and exhibiting fluid inclusions with homogenization temperatures around ~100°C, thereby possibly suggesting a possibly advanced burial stage or an abnormal geothermal gradient linked to volcanic activity (Girard & San Miguel, 2017; Girard et al. 2017).

5.3. Insights from the partition between length-slow and length-fast fibrous microquartz

Petrographic and geochemical investigations of the quartz cements highlights a bimodal partitioning amongst the different studied sites, based on: (1) the length-fast vs. the length-slow habit of the fibrous microquartz, (2) the greater carbonate dissolutions associated (prior) to length-fast fibrous microquartz precipitation, and (3) the increase in δ¹⁸O_{quartz} values from length-fast to length-slow fibrous microquartz. We suggest that this textural and geochemical
partition between the two fibrous microquartz habit may be an indicator of the environmental conditions of precipitation. Originally, Folk & Pittman (1971) described these quartz habit. Warren (2016) reviewed and discussed the latter in light of their precipitation processes. Length-fast cements would be associated with neutral to acidic environments because high Si concentrations and low pH favor the polymerization of monosilicic acid (Si(OH)_4) groups. Such groups would likely precipitate tangentially on the precipitation surface (Folk & Pittman, 1971). In contrast, high pH tends to favor the occurrence of monomeric Si(OH)_4 tetrahedra, inducing that length-slow fibrous microquartz reflect alkaline environments. The chemical behavior of silica in function of pH is explained by Iller (1971). Arbey (1980) discussed extensively Folk & Pittman’s (1971) observations and pointed out the potential control of dissolved sulphates on quartz precipitation. Sulphate-rich environments would favor the precipitation of length-slow fibrous microquartz cements, whereas sulphate-poor environments would favor length-fast forms. Unfortunately, petrographic or geochemical evidence for a former presence of sulphates in the studied samples is lacking. Bustillo et al. (2012) reported similar observations as Folk & Pittman (1971) in the Miocene continental cherts from the Madrid Basin (Spain), and linked the occurrence of length-fast fibrous forms to carbonate sedimentary settings lacking dolomite or gypsum. Conversely, they related the precipitation of length-slow fibrous forms to Mg-rich palustrine evaporitic environments associated with gypsum, dolomite, and Mg-clay minerals. As explained previously, the most probable Si source leading to quartz precipitation in the presalt series is alteration of volcanic rocks, with a possible contribution of mantle rocks through hydrothermal activity. Deep hydrothermal contribution in rift settings linked to the mantle has reported by Lee et al. (2016). The alkalinity, pH, and elemental concentrations of the presalt diagenetic fluids may have been strongly controlled by the nature of rocks prevailing in the palaeohydrological catchment/basin (e.g. Teboul et al., 2017). Teboul et al. (2016) documented such a process in
recent and modern continental carbonate settings: fluid/rock interactions involving mantle rocks in the Voltri massif of the Ligurian ophiolites (Italy) lead to highly alkaline fluids, in agreement with other observations on Oman Ophiolites (Chavagnac et al., 2013; Leleu et al., 2016). Similarly, it has been demonstrated that fluid/rock interactions of basic volcanic rocks of Reunion Island (Indian Ocean) lead to neutral to alkaline fluids (Teboul et al., 2016). The alteration of presalt acid volcanic rocks in the offshore Kwanza basin (Angola) would have led to near neutral to slightly acid fluids, depending on the amount of CO₂ during fluid/rock interaction (Teboul et al., 2017). Deocampo & Renaut (2016) emphasized such interaction in the soda lake of the east African rift. As shown by several authors (Pereira, 1969; 1971; Marzoli et al., 1999; Comin-Chiaramonti et al., 2011), the early Cretaceous volcanic rocks from Angola include a wide spectrum of compositions, from basalts to rhyolites, while that of late Cretaceous volcanic rocks ranges from basalts to trachytes. In light of these results, we assume that variable water chemistry and pH could have occurred in presalt palaeohydrological basins, reflecting the heterogeneity of presalt basement and strata, and structural heritage. We therefore postulate two major end members. The first implies ultramafic/basic volcanic rocks, which can lead, through fluid/rock interaction, to the production of Si-rich alkaline diagenetic fluids. Such fluids would be prone to form length-slow fibrous microquartz. The second end member results from interactions with intermediate/acid volcanic rocks, producing Si-rich neutral to acidic diagenetic fluids, prone to form length-fast fibrous microquartz. The occurrence of significant dissolution features observed prior to silica cementation supports the hypothesis of an acidic diagenetic fluid associated with length-fast fibrous microquartz, in Mina do Gesso, Mariquita, and Toca-1 studied sites.

From a thermodynamic point of view, the solubility of silica polymorphs is significantly greater in strongly basic pH conditions (e.g. greater than 9 at 25°C or greater than 8 at 100°C;
Fig. 7; Iler, 1979). This implies that under basic conditions supersaturation will be reached only at extremely high dissolved silica concentrations. Fig. 7 also illustrates that temperature plays a prominent role in the precipitation of silica polymorphs. It can be seen that the solubility of silica polymorphs is about one order of magnitude greater at 100°C compared to 25°C. Consequently, high temperature waters are prone to carry high dissolved silica content and any decrease in fluid temperature would induce silica precipitation, either as fibrous microquartz or quartz depending on the initial dissolved silica concentration (Fig. 7).

The microporous fabric of the length-slow fibrous microquartz observed in Shrubby-1 is a peculiar texture/feature. To our knowledge, only a few authors described similar cements in deeply buried carbonate sequences, e.g. Saller et al. (2016) in the Kwanza basin (Angola) and Watney & Rush (2012) in the Arbuckle Group saline aquifer (Kansas, USA). In the present study, observations indicate that the porous fibrous microquartz shows a strong interrelationship with oil, suggesting that oil emplacement may have started coevally or shortly after fibrous microquartz precipitation. The decomposition of organic matter may also favor early silica precipitation as increasing CO₂ partial pressure, through oxidation of organic matter, leads to a simultaneous increase of carbonate solubility and decrease of silica solubility (Gao & Land, 1991; Smrzka et al., 2015). It is commonly accepted that fibrous microquartz probably indicates extremely high precipitation rate in highly supersaturated fluids (García-Ruiz, 2000; Warren, 2016). The migration of hydrocarbons soon after precipitation of microquartz fibers may have inhibited further development of quartz overgrowths on the fibrous microquartz, thus preserving its microporosity. An early oil migration in a rift setting can result from the prevalence of elevated thermal regimes, possibly associated to volcanic activity (Ziegler, 1992; Pindell, 1995; Chen et al., 1999). The Limagne Basin (Massif Central, France) represents a valuable analog of a continental rift system showing the relationship between magmatic activity, carbonate precipitation, oil migration,
and siliceous cementation in shallow buried lacustrine sediments (Wattinne, 2004; Couturié, 2015).

It has been noticed above that high $\delta^{18}O_{\text{quartz}}$ occurs in porous length-slow microquartz cements in Toca-2 and Shrubby-1 compared to the length-fast non-porous microquartz found in Toca-1, Berilo and Mina do Gesso (Fig. 6). In former studies, $\delta^{18}O_{\text{quartz}}$ measurements on siliceous cements have mainly been carried out on cements formed from seawater in Precambrian-Phanerozoic marine sediments with the aim of investigating their temperature of formation (Knauth & Lowe, 1978; Marin-Carbonne et al., 2014; Knauth & Epstein, 1976). In such studies, reconstructing the temperature of formation of the silica cement based on its $\delta^{18}O_{\text{quartz}}$ signature is possible because they formed from seawater of which the oxygen isotope composition can be constrained. Such an approach cannot be used for the early silica cements of the presalt deposits, because the parent water is thought to be of continental origin and its oxygen isotope value is unknown. Microthermometry measurements of primary fluid inclusions in the silica cement could help constraining their formation temperature. However, the presence of fluid inclusions is impossible to ascertain due to the fibrous nature of the cement. Nevertheless, indirect temperature and isotopic constraints can be derived from the published information. According to palaeoenvironmental reconstructions proposed by Chaboureau et al. (2013), the studied area was located in the tropical zone during Aptian time (Fig. 1B). This suggests that lake water temperature was unlikely to be much below ca. 25°C. Accordingly, we used this value as a minimum temperature of crystallization for early fibrous microquartz cements in Fig. 8A (“T”). According to Shackleton & Kennett (1975) and Sheppard (1986), the mean isotopic composition of the ocean before the Miocene was about $\delta^{18}O_{\text{water}}$ = -1‰. However, Veizer et al. (1986) showed that the seawater composition was not uniform through time. Although, the same author showed that the major variation occurs in the Early and Mid-Paleozoic oceans. Considering that the $\delta^{18}O_{\text{water}}$ of Aptian sea water was
on the order of -1‰, all \( \delta^{18}O_{\text{water}} \) values below -1‰ can be interpreted as tracing a meteoric influence, and values greater than 0‰ can be attributed to magmatic, evaporated or highly modified waters (Criss, 1999; Hoefs, 2009). Magmatic waters are defined by Criss (1999) as waters of any origin that has equilibrated with magma. Following Sheppard (1986), the \( \delta^{18}O_{\text{water}} \) values of “primary magmatic water” can reach approximately 10‰, following equilibrium with crystalline rocks at igneous temperatures. Similar isotopic composition would be expected in a volcanic setting where deep hypogean fluids are exposed to igneous temperatures (Sheppard, 1986; Taran et al, 1995). These waters differ from hydrothermal waters, which are predominantly derived from local meteoric waters (Craig et al., 1956), and can show elevated \( \delta^{18}O_{\text{water}} \) values (up to 20‰ or more) as a result of water/rock interactions with \(^{18}\)O-enriched rocks. Of course, evaporation processes can also produce significantly \(^{18}\)O-enriched waters (up to 10‰). High to extremely high \( \delta^{18}O_{\text{water}} \) values have been reported in evaporative settings, i.e. 11.3‰ in Lake Baringo, Kenya (Renaut et al., 2002) or 31.3‰ in small, shallow lakes of the Western Sahara (Gonfiandini, 1986), but such values remain exceptional. Boiling can also enhance \(^{18}\)O\(_{\text{water}}\)-enrichment (Bottinga & Craig, 1969). This shows that during boiling, \(^{18}\)O partitioning into the liquid phase can lead to strongly enriched \(^{18}\)O\(_{\text{water}}\). The above information and constraints provide an interpretative frame for our results, which are further discussed below.

The large spread in \( \delta^{18}O_{\text{quartz}} \) data reported in this study is noteworthy (Fig. 6). This heterogeneity of isotopic values can reflect temperature variations or changes in fluid composition at microsite scale. Processes other than temperature and initial parent \( \delta^{18}O_{\text{water}} \) may affect the \( \delta^{18}O \) of quartz/silica; including growth under disequilibrium conditions (Dickson, 1991), sector zoning (Jourdan et al., 2009), variable incorporation of trace Al in the quartz lattice due to a change from normal step-wise growth to spiral growth (Yoshimura et al., 1979; Larkin et al., 1982). In fibrous microquartz cements, minor \( \delta^{18}O \) variations could
also be related to the occurrence of OH-related impurities. In general, fibrous microquartz cements tend to contain approximately 2 wt.% of silanol (SiOH) and water (H\textsubscript{2}O), with water located mainly in very small fluid inclusions (Flörke \textit{et al.}, 1982; Schmidt \textit{et al.}, 2013). The strong variation of $\delta^{18}$O values observed by using SIMS analysis highlight the necessity to be careful of bulk quartz and chert $\delta^{18}$O as it can homogenize true values of multiple quartz generations. Similar observations have been shown in several recent studies (Denny \textit{et al.}, 2017; Cammack \textit{et al.}, 2018). However, the possible influence of the processes listed previously in the large $\delta^{18}$O values spread observed in this study cannot be assessed at this time. As a consequence, it will not be discussed further.

Despite the difficulty to use $\delta^{18}$O\textsubscript{quartz} as a geothermometer, the elevated $\delta^{18}$O\textsubscript{quartz} values measured in the Presalt cements provide valuable information. Saller \textit{et al.} (2016) suggested that such elevated values reflect an evaporitic source of the quartz-forming waters in the presalt sediments. In our study, the oxygen isotopic compositions of water have been extrapolated to low temperatures based on the fractionation factor of Matsuhisa \textit{et al.} (1979) for the quartz-water system. We consider that there is no significant differences between this fractionation factor and the calibrations of Clayton \textit{et al.} (1972) or Hu & Clayton (2003). The length-slow porous microquartz cements show a maximum value of 37‰ (Fig. 6). Considering the minimum temperature of $T=25^\circ$C, the $\delta^{18}$O\textsubscript{water} values would be of at least 0‰ (Fig. 8A) and possibly higher if a higher temperature of formation is considered, i.e. 15‰ at a temperature of 100 °C (Fig. 8A) which would represent extreme conditions of water evaporation. High $\delta^{18}$O\textsubscript{quartz} (44‰) were also obtained from microquartz precipitating in evaporative environments from the East African Rift (Fig. 9; O’Neil & Hay, 1973). Literature data of quartz precipitated during early to late marine diagenesis exhibit $\delta^{18}$O\textsubscript{quartz} values consistent with the global range compiled in this study (Fig. 9). In the past, several authors interpreted high isotopic values as the expression of very low temperature of
crystallization (usually <5 °C), consistent with deep oceans conditions (Degens & Epstein, 1962; Knauth & Epstein, 1975). As indicated above, such low temperatures are incompatible with the presalt depositional environments. As shown in Fig. 9, the range of δ\textsuperscript{18}O\textsubscript{quartz} values (20-40‰) measured in the Presalt silica cements has never been documented in any other non-evaporitic sedimentological or diagenetic setting.

The length-fast and length-slow non-porous microquartz cements exhibit statistically lower δ\textsuperscript{18}O values (27-32‰) than their length-slow porous counterpart (33-37 ‰). The lowest value for length-fast fibrous microquartz in Toca-2 is 27‰, which implies that the δ\textsuperscript{18}O\textsubscript{water} would have been lower than 0‰ (i.e. meteoric or mixed meteoric-marine) at all temperatures below 60 °C (Fig. 8A). For the megaquartz cements, the highest δ\textsuperscript{18}O\textsubscript{quartz} of 40‰ was measured in Shrubby-1 (Fig. 6). Saller \textit{et al.} (2016) reported a similar maximum δ\textsuperscript{18}O\textsubscript{quartz} value of 41‰ in the presalt megaquartz cements in offshore Kwanza (Fig. 9). Girard & San Miguel (2017) and Girard \textit{et al.} (2017) reported primary two-phase fluid inclusions in megaquartz cement in the Presalt carbonates of an offshore Kwanza well yielding homogenization temperatures clustering around ~100°C. The latter authors also emphasized that the presalt deposits at that location were submitted to elevated temperatures, approaching ~200°C. Poros \textit{et al.} (2017) documented homogenization temperatures ranging from 98 to 123°C for fluid inclusions occurring in chalcedony and megaquartz cements in the Omosi-1 well, offshore Kwanza. Based on this information we can constrain the temperature of crystallization of the megaquartz cement in our samples to be between 100 and 200°C (Fig. 8B, “T2” and “T3”). At such temperature, the measured δ\textsuperscript{18}O\textsubscript{quartz} values would mainly reflect precipitation from highly \textsuperscript{18}O-enriched waters. Only the two lowest δ\textsuperscript{18}O\textsubscript{quartz} values in our dataset would be compatible with a modified marine water signature (Fig. 8B). Girard \textit{et al.} (2017) favor the influence of pervasive hydrothermal waters based, among other things, on the omnipresence
of highly saline fluid inclusions (>200 g/l) in the successive cements (dolomite, calcite, quartz) found in the studied samples of the presalt carbonates.

5.4. Proxies controlling silica precipitation: scenarios

Following the elements discussed above, we propose that the different habit and geochemistry of the different quartz cements were probably controlled due to fluid/rock interaction with two substratum end-members (respectively intermediate to acid substratum and basic to ultramafic substratum). From basin to pore scale, four main steps from early to late burial diagenesis can then be described as follows (Fig. 10):

- Step 1: at basin scale, at several sites (Mariquita, Berilo, Toca-1, Mina do Gesso), 18O-depleted fresh to slightly evaporated waters could have interacted with intermediate to acid volcanic substrates, promoting circulation of Si-rich neutral to acid waters in the sediment (Fig. 10-1A). At other sites (Shrubby-1, Toca-2), hypogean fluid/rock interaction between 18O-enriched slightly to highly evaporated water and basic to ultramafic basement would promote intra-sedimentary circulation of Si-rich alkaline waters (Fig. 10-1B). In both case, the initial fluid isotopic composition could have been undifferentiated if we consider that the final 18O composition of the water is only influenced by water/rock interaction along the hydrogeological pathway.

- Step 2: at pore scale, after precipitation of carbonate cements close to the sediment/water interface, circulation of Si-rich waters during early diagenesis induced partial carbonate dissolution, especially when Si-rich acidic waters interacted with the sedimentary body (Fig. 10-2A).

- Step 3: in agreement with Folk & Pittman (1971), the intense circulation of neutral to acidic waters, very rich in Si(OH)₄ polymers (due to acidic pH), promoted rapid
precipitation of length-fast fibrous microquartz displaying the lowest $\delta^{18}$O$_{\text{quartz}}$ values (Fig. 10-2A to 3A). Conversely, the intense circulation of alkaline waters very rich in ionized Si(OH)$_4$ tetrahedra promoted rapid precipitation of length-slow fibrous microquartz characterized by the highest $\delta^{18}$O$_{\text{quartz}}$ values (Fig. 10-2B to 3B). In case of early oil impregnation, the length-slow fibrous microquartz cement may have preserved microporosity between individual fibers during later diagenetic processes (Fig. 10-3B').

- Step 4: at most sites, the silica diagenesis ended with megaquartz precipitation during early to burial diagenesis, usually associated with hydrothermal circulations (Girard & San Miguel, 2017; Girard et al., 2017; Poros et al., 2017) possibly driven by geothermal convection below the salt cover (Jones & Xiao, 2013). Highly $^{18}$O-rich megaquartz cementations followed the precipitation of length-slow fibrous microquartz (Fig. 10-4B), in accordance with the interaction of strongly enriched waters along this diagenetic pathway. Further work is needed to better constrain the conditions of precipitation of the late quartz cements.

6. Synthesis and conclusions

The integrated petrographic and geochemical study of quartz cements occurring in the presalt rocks from offshore and onshore basins of the West African margin provides valuable information to constrain the palaeoenvironmental and diagenetic settings. This study is the first attempt to combine $\delta^{18}$O$_{\text{quartz}}$ values with the length-fast vs. length-slow habit of fibrous microquartz cements. Several potentially useful markers are highlighted by the study and are summarized below:

- Early length-slow fibrous microquartz cements precipitated from Si-rich, alkaline waters. $\delta^{18}$O$_{\text{quartz}}$ values range from 31 to 37‰e, reflecting formation from evaporated
water at probably low temperature (<100°C). The required alkaline nature of these waters is probably linked to fluid/rock interactions involving basic volcanic rocks, with a possible mantle influence. A mantel influence would be particularly consistent with the recent geodynamic models established for the South Atlantic margins. Early to late length-fast fibrous microquartz cements with $\delta^{18}O_{\text{quartz}}$ values ranging from 27 to 32‰, precipitated from moderately to non-evaporated water at higher temperatures. Influence of marine water cannot be excluded if temperature was between 35 and 60°C. The length-fast cements require Si-rich neutral to acidic waters, probably derived from interaction with intermediate to acid volcanic rocks. Such interactions are consistent with field and well data, documenting the existence of magmatic intrusions and hydrothermal alteration. The characterization of length-fast vs. length-slow habit of fibrous microquartz holds great potential to unravel the paleo pH history of the parent waters.

- A peculiar form of length-slow fibrous microquartz cements is described which shows abundant preserved interfibre porosity often impregnated by oil. Such cements have not been investigated in the literature and would certainly deserve more attention in order to unravel the link between early oil migration and early quartz precipitation in alkaline, saline waters.

- Megaquartz following length-slow fibrous microquartz exhibits $\delta^{18}O_{\text{quartz}}$ values up to 40‰, that are indicative of highly $^{18}$O-rich waters, most likely including a significant contribution of recycled evaporated waters or strong water/rock interaction. Strong variations in $\delta^{18}O_{\text{quartz}}$ values can reflect temperature variations or changes in fluid composition. This would indicate a strong heterogeneity of thermo-chemical conditions at microsite scale. Megaquartz is likely associated with lower cementation
rate and dissolved silica concentration in the diagenetic waters compared to fibrous microquartz.

In order to further investigate and understand the conditions of quartz cementations in the studied Presalt deposits from the South Atlantic more detailed fluid inclusion work should be conducted. Such an approach should also be applied to the onshore and offshore Presalt basins of the Brazilian margin, in order to reconstruct a global and possibly predictive model of quartz cementation in Presalt series. Moreover, several authors have highlighted that dissolved sulphates may also control the habit of quartz during precipitation. Unfortunately, no information on the occurrence of former sulphate-rich minerals was recorded in the studied intervals.

Acknowledgments

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Figure captions

**Figure 1:** (A) Location map of the studied wells and outcrops in the Congo, Kwanza, and Namibe basins (modified from Torsvik, *et al.*, 2009; Matthews *et al.*, 2011; Heine *et al.*, 2013; Seton *et al.*, 2014; Péron-Pinvidic *et al.*, 2015). (B) Palaeogeographical map of the presalt basins at the Lower Aptian (124-123 Ma) replacing the studied material in lacustrine environments (modified from Chaboureau *et al.*, 2013).

**Figure 2:** Schematic representation of typical diagenetic sequences (Not to scale; AC: aragonite to calcite neomorphism; Ank: ankerite; BC: blocky calcite; Cal.: calcite; Dol.: dolomite; F. BC: ferroan blocky calcite; FC to dol.: radiofibrous calcite to dolomite; FQ: fibrous microquartz; pFQ: porous fibrous microquartz; LQ: laminated microquartz; MQ: megaquartz; Ox.: oxides; Sdl.: saddle dolomite; Sdl. to Cal: saddle dolomite replaced by calcite). Major fracturing and dissolution stages are reported as f(n) and d(n) respectively. Colours of carbonates and sulphates refer to cathodoluminescence observations. Quartz cements are in blue. Notice that time is not constrained, all sequences can be considered as asynchronous.

**Figure 3:** Photomicrographs of Mariquita (A-E) and Mina do Gesso (F-G), onshore Kwanza Basin. (A) Cross-polarized light (XPL) view of fibrous microquartz cement (FQ) followed by megaquartz (MQ) in a calcitic matrix (Cal.) of microbial origin. (B) XPL view of botryoids of radiaxial calcite, which has been successively dolomitized (Dol.), fractured and enclosed in a megaquartz cement (MQ). (C) Plain polarized light (PLL) view of fibrous microquartz cement (FQ) followed by megaquartz (MQ). (D) XPL view of a dolomitized laminated crystalline crust that underwent fracturing and dissolution before megaquartz precipitation.
(adapted from Foubert et al., 2014). (E) Location of some of the SIMS measurements in the megaquartz of Mariquita. Note the general $^{18}$O enrichment through time. (F) XPL view of partial fibrous microquartz precipitation along several planes of a calcitic (Cal) spherulite (yellow arrows). (G) PPL (left) and XPL (right) of fibrous microquartz cement in the fenestral porosity of micritic microbial mats.

**Figure 4:** Photomicrograph and SEM pictures of Shrubby-1 (Chela Fm.), offshore Kwanza Basin. (A) PPL (left) and cathodoluminescence (CL; right) views of quartz precipitations in pores. A first generation porous fibrous microquartz (pFQ1) is followed by a thin isopachous layer of laminated microquartz cement (LQ1), and by megaquartz cement (MQ). The initial shrubs that constitute the substratum underwent several stages of dolomitization (Dol) and dissolution. (B) Close-up view of pFQ1, LQ1, and MQ under UV epifluorescence highlighting the porous nature of pFQ1 and LQ1 (blue fluorescence due to the impregnated epoxy resin). (C) Close-up view of pFQ1, LQ1, and MQ under PPL. Some patches of microquartz (µQ) occur. Notice the bitumen occurring sporadically inside the porosity of pFQ1. (D) PPL view highlighting pFQ1 and pFQ2. Notice the exceptional porosity of pFQ1 (blue due to the epoxy resin). pFQ2 has a petaloid texture and pores are filled by bitumen. (E) PPL view of length-slow fibrous microquartz determined using a first order (full wave) retardation plate. (F) SEM view of laminated microquartz cement (LQ) followed by megaquartz cements (MQ). (G) SEM view of aligned anhedral to euhedral microquartz crystals along fibers and planes of the pFQ1 fibrous cement (yellow arrows). (H) SEM view of megaquartz cement (MQ). (I) SEM element map highlighting the porous nature (dark grey) of pFQ1 and LQ1.
Figure 5: Photomicrographs of Shrubby-1 (Cuvo Fm., offshore Kwanza; A-B), Berilo (offshore Kwanza; C), Toca-1 (offshore Congo; D-E), and Toca-2 (offshore Congo; F-G). (A) PPL view of multiple megaquartz (MQ1 and MQ2) and porous laminated microquartz (LQ1) cements occurring in a fracture. The remaining porosity is partially filled by some blocky calcite (BC). (B) PPL view of LQ1, MQ1, and BC cements occurring in a fracture. (C) PPL view of brownish fibrous microquartz (FQ) and megaquartz (MQ) cements in fenestral porosity. (D) PPL view of fibrous microquartz cements (FQ) infilling primary porosity in the coquina facies of Toca-1. The pelecypod shells, underlined by yellow dashed lines, underwent dissolution of aragonite to calcite neomorphism. (E) Localization of SIMS $\delta^{18}$O$_{\text{quartz}}$ measurements in the FQ cements of Toca-1 under PPL view. (F and G) PPL (left) and CL (right) views of FQ and MQ cements infilling the porosity in the coquina facies of Toca-2. The yellow dashed lines underline the recrystallized and dissolved bivalve shells.

Figure 6: SIMS $\delta^{18}$O$_{\text{quartz}}$ (‰ SMOW) according to textural habit of quartz cements in the six studied sites. Colours of length-fast and non porous length-slow fibrous microquartz were determined according to microscopic observations using a $\lambda$-retardation plate. Colour of the microporous length-slow fibrous microquartz according to the blue epoxy resin observed in the porosity.

Figure 7: Silica solubility curves as a function of pH at 25 and 100°C, calculated using PHREEQC. Thermodynamic data come from the Thermoddem database (http://Thermoddem.brgm.fr/; Blanc et al., 2012). Log K are given values at standard state conditions (T = 298 K, P = 1 atm). The log K of chalcedony has been considered as equivalent of fibrous microquartz. Log $K_{\text{quartz(alpha)}} = -3.740$; Log $K_{\text{chalcedony}} = -3.456$. 
Figure 8: $\delta^{18}$O$_{\text{quartz}}$ (‰ SMOW) as a function of temperature (°C) for (A) the fibrous microquartz cements and (B) the megaquartz cements. Oxygen isotopic compositions of water (dashed lines) were calculated using the quartz-water fractionation equation of Matsuhisa et al. (1979): \[1000 \ln \alpha_{\text{quartz-H}_2\text{O}} = 3.34 \times (10^6 T^{-2}) - 3.31.\] T, T1, T2 and T3 refer to the hypothetical minimum and maximum temperatures of crystallization (cf. discussion).

Figure 9: Range of $\delta^{18}$O$_{\text{quartz}}$ (‰ SMOW) values in the present study and comparison with values derived from a literature review of $\delta^{18}$O$_{\text{quartz}}$ in carbonate rocks, according to quartz morphologies and precipitation settings (A: Ewers, 1991; B: Sharp et al., 2016; C: Clayton & Steiner, 1975; D: Herzig et al.; 1988; E: Shanks et al., 2007; F: Renaut et al., 2002; G: Saller et al., 2016; H: O’Neil & Hay, 1973; I: Degens & Epstein, 1962; J: Kolodny & Epstein, 1976; K: Knauth & Epstein, 1975; L: McBride et al., 1999; M: Harris, 1989; N: Abruzzese et al., 2005; O: Jourdan et al., 2009). Precipitation settings are defined according to the published interpretations. Laboratory techniques used for $\delta^{18}$O$_{\text{quartz}}$ measurements (bulk or microsampling, laser fluorination or SIMS) are specified.

Figure 10: Relative timing and proxies controlling silica cements in the presalt deposits of the West African margin, according to their petrographic and isotopic evolution. Please refer to the part 5.4 of the discussion for explanations. Colours in 3A and 3B are representative of microscope observations with a $\lambda$-retardation plate. The colour in 3B’ is representative of the blue epoxy resin impregnating the microporosity of those fibrous microquartz.
Onshore Namibe Basin (Angola)

Offshore Kwanza Basin (Angola)

Lower Congo Basin (Congo)
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**Laminated microcrystalline**

- **Zone 1**
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  - Site B: X X X

- **Zone 2**
  - Site A: X X X

**Lamprophyres**

- **Zone 1**
  - Site A: X X X
  - Site B: X X X

**Metamorphic Rock**

- **Zone 1**
  - Site A: X X X
  - Site B: X X X

**Nappes**

- **Zone 1**
  - Site A: X X X
  - Site B: X X X

**Phaneritic Sill**

- **Zone 1**
  - Site A: X X X
  - Site B: X X X

**Phaneritic Blende**

- **Zone 1**
  - Site A: X X X
  - Site B: X X X
A  Fibrous microquartz

B  Megaquartz

T°C

δ¹⁸O

δ¹⁸O_pore (% SMOW)

δ¹⁸O_parc (% SMOW)
Quartz cements show extremely high $\delta^{18}O$ values (up to 40‰)
- In situ $\delta^{18}O$ analyses reveal a variability between and within different quartz generations
- $\delta^{18}O$ associated with length fast and length slow quartz habits helps to understand paleofluid history.