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Probing multi-scale structure and dynamics of waxy crude oil by low field NMR, X-Ray scattering and optical microscopy

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crystallization, bulk

ABSTRACT

Wax deposition is one of the major concern for waxy crude oil production and transportation. A better understanding and prediction of fluid properties related to this issue requires knowledge of the medium structuration at scales ranging from nanometers (molecules) to a few micrometers (crystals). For this purpose, the behavior of a waxy crude oil in bulk was compared to a model oil over a wide range of temperatures above and below the Wax Appearance Temperature (WAT). The combined use of Cross Polarized Microscopy (CPM) and the implementation of innovative techniques for the field such as SAXS/WAXS and low field NMR has provided a more precise idea of the structure of these two types of fluids. If the nature of the orthorhombic crystals and their lamellar shape are identical for both fluids, a very appreciable difference is highlighted in their chain axis dimensions. The *n*-paraffins crystals size is larger than 120 nm in the model oil. In the crude oil, it is only about 1-10 nm with a long range order in the directions perpendicular to the chain axis and a monomolecular thickness in the chain axis. Moreover, from the model oil CPM images, we observed aggregates of lamellar shape crystals. Since the model oil does not contain asphaltenes and resins, the crystals are larger and the branches divide significantly less than in the crude oil which results in a trapped liquid protons population unobservable in the model oil. All these observations give a vision of the structure of *n*-paraffins crystals. It is made of aggregates of relatively dense lamellae in the center and more aerated lamellae at the periphery which split into several branches.

1. INTRODUCTION

Production and transportation of waxy crude oils through long pipelines at low temperatures is a critical challenge in deep and ultra-deep offshore. Waxy crude oils are complex mixtures containing paraffins, aromatics, naphthenes, resins and asphaltenes with a carbon number higher than 20¹. At high temperatures, the *n*-paraffins (waxes) remain dissolved in crude oil but when the temperature at the pipe wall drops below the Wax Appearance Temperature (WAT), the crude oil *n*-paraffins start to crystallize and cause serious issues such as wax deposition on the walls of pipelines²⁻⁴, an increase of the waxy crude oil viscosity leading to a non-Newtonian behavior⁵⁻⁷ and restart issues of the pipelines due to the waxy gelation during shutdowns⁸⁻¹⁰. These problems lead to significant additional production costs¹¹ and the development of operational remediation techniques such as pigging or chemical inhibition requires a deep understanding of the complex behavior of waxy crude oils. In order to reproduce the flow of crude oil in pipelines, flow loop experiments are carried out in the laboratory and require the knowledge of the properties of *n*-paraffins crystals aggregates such as structure, size of unit crystals, fluid behavior in aggregates ...). Indeed, knowing these properties enable a better characterization of wax deposit structure and a deeper understanding of the different mechanisms that lead to wax deposit formation.

The literature presents numerous studies performed in order to probe the structure of the deposit in which classical characterization techniques have been used. These techniques include rheology^{11,12,13}, DSC^{11,12,14}, HTGC^{11,15-17} and densimetry⁶. They allow the determination of the waxy oils main properties such as viscosity, density, WAT, Pour Point, distribution and the *n*-paraffins content. These fundamental characterizations are important for the development of

models predicting wax deposit formation and flow properties but do not allow a multi-scale description of the deposit structure.

To investigate the morphology and size of crystals in crude oils, optical microscopy has been widely used but, depending on the experimental conditions set-up, different sizes and morphologies have been found^{14,18–20}.

Slightly more advanced techniques have been implemented for waxy oils characterization. NMR relaxometry has been used to characterize waxy crude oils and in particular to determine the *n*-paraffins content in crude oils and their distributions^{21,22}. Furthermore, NMR relaxometry has also been used to understand the role of inhibitors on wax crystallization²³. X-ray diffraction has been used to study the structural behavior of pure *n*-paraffins or model oils, but few studies have been carried out directly on the crude oils^{24–26}. Indeed, binary, ternary, quaternary mixtures or model oils have been used mainly because of their simplified structure compared to real crude oils. However, these studies are still far from real complex systems.

In the present paper, we propose to make a description of the structure of a waxy crude oil at multi length-scale (1 Å to 10 μm) in both model oil and crude oil using a variety of analytical techniques such as Cross Polarized Microscopy (CPM)^{19,27–29}, Small and Wide angle X-ray scattering (SAXS/WAXS)^{30–32} and low field NMR^{33,34}. Each technique provides useful information to finely describe the structural behavior of *n*-paraffins crystals in a model and a crude oil: CPM makes the visualization of the *n*-paraffins crystals possible, SAXS/WAXS provides information on crystals inter-arrangement, their size and shape and NMR can be used to study fluid dynamics within the formed crystal network.

2. MATERIALS AND METHODS

In this study, a model oil and a crude oil were used. The model oil allowed to simplify the application of CPM and to evaluate its ability to represent a crude oil according to the observed behaviors.

2.1. Crude oil and model oil

The model oil was prepared using a disaromatized aliphatic oil (Hydrosel G250H (C₁₅-C₂₀), number C.A.S 64742-46-7, crystallization temperature -20 °C) density 0.812 g/cm³ at 20 °C and 10 % weight of wax (commercial wax purchased from Prolabo, melting point 52-54 °C). The model oil WAT is 22°C, which is well suited to the different experimental set-ups we used. The waxy crude oil, which was provided by Total and coming from a Congo field, has a density of 35° API (American Petroleum Institute gravity) and a WAT of 51 °C. The crude oil was filtrated with a filter size of 5 µm at 60 °C in order to get rid of all solid particles such as sand particles.

Before the beginning of each experiment, samples were heated at a temperature above the WAT (WAT+20°C) during few hours and were hand-mixed during approximately 1 minute.

The WAT and the weight fraction of wax crystals C_{wax} (wt.%) of the oils were obtained with a differential scanning calorimeter (DSC). The detailed procedure is described in supplementary information **a**. Volume fractions ϕ (where $\phi = V_s / (V_s + V_L)$ with V_s as being the solid volume and V_L the liquid volume) were obtained from C_{wax} and densities of the suspending liquid phase ρ_L (supplementary information **b**) and solid wax ρ_{wax} . According to the literature, between nC_{20} and nC_{32} , the density of the solid phase varies slightly and linearly between³¹ 0.92 g/cm³ and 0.93 g/cm³. Since it is known that a n -paraffins mixture produces a solid solution that contains

defects, then the density of the mixture will be lower than that of the pure phase of *n*-paraffins which constitutes the mixture³⁵. Thus, we assumed that the density of the solid wax in the mixtures^{6,36} ρ_{wax} is about 0.9 g/cm³.

A SARA analysis was performed on the crude oil in order to know its composition (**Table 1**).

Table 1. *Composition of the crude oil.*

Components	Content, % mass
C ₆ -C ₁₄	25.8
Saturated C ₁₄₊	49.7
Aromatics C ₁₄₊	7.6
Resins	13.2
Asphaltenes	3.6

The *n*-paraffins distribution of the crude oil and the commercial wax are shown in **Figure 1**. The *n*-paraffins distribution of the commercial wax was obtained by High Temperature Gas Chromatography (HTGC) with an Agilent 6890 gas chromatograph. HTGC is commonly used to determine the *n*-paraffins distributions in crude oils and its products^{15,16,37}. The *n*-paraffins distribution in the crude oil was obtained by performing GC analysis with a Thermo Scientific Trace 1300 gas chromatograph on the fraction which does not contain heavy resins and asphaltenes (obtained by SARA separation). The *n*-paraffins distribution of the commercial wax covers 13 *n*-alkanes ranging from C₂₁ to C₃₄ and carbon distribution of the crude oil had a range from C₇ to C₄₀₊. The *n*-paraffins proportion was quantified from C₇ to C₄₀ and it is approximately 26 weight %. A simulated distillation analysis by HTGC was also carried out on the crude oil and showed that the distribution of *n*-paraffins extends to over 86 carbon atoms. However, it was not possible to quantify them. The *n*-paraffins distribution is characterized by its mean number of

carbon atoms $\bar{n}$ and the standard deviation σ . These parameters were calculated from relationships^{26,38} described in supplementary information **d**.

The characteristics of the *n*-paraffins distributions in the model oil and crude oil (only from C₇ to C₄₀) are summarized in **Table 2**. For the crude oil, since we do not have the complete distribution we could not determine $\bar{n}$ and we can only calculate a minimum value of σ .

Table 2. *Characteristics of the crude oil and model oil n-paraffins distributions.*

Sample	Composition in <i>n</i> -paraffins	<i>n</i> -paraffins content	$\bar{n}$	σ
Crude oil	C ₇ H ₁₆ to C ₄₀ H ₈₂ ⁺	> 26 wt. %	-	>7.2
Model oil	C ₂₁ H ₄₄ to C ₃₄ H ₇₀	10 wt. %	24.8	2.1

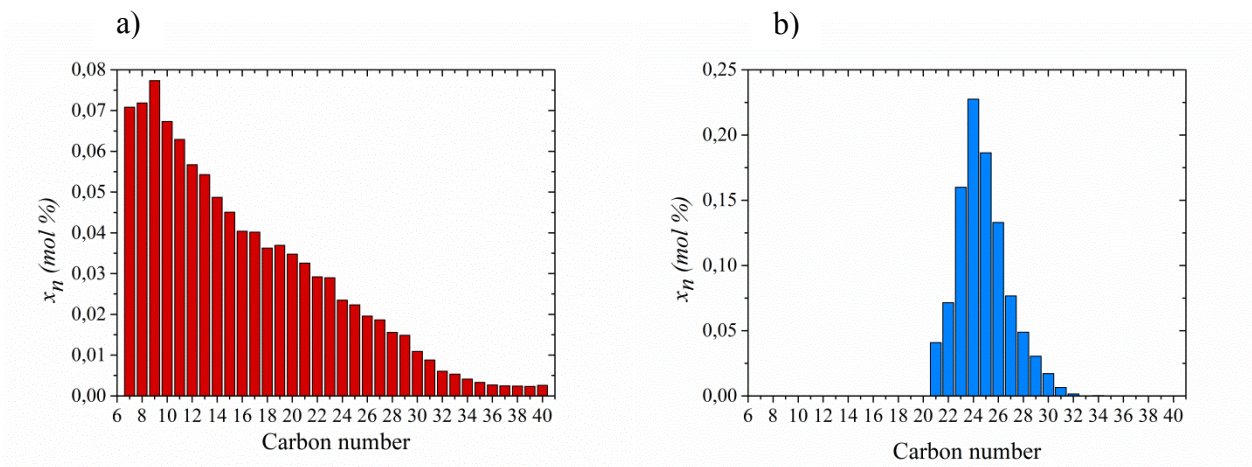


Figure 1. *n*-paraffins molar distributions in a) the crude oil (from C₇ to C₄₀) and b) the model oil.

According to the GC, the minimum mass of paraffin contained in the crude oil is 26 weight %. We considered this value to be the total mass of *n*-paraffins in the sample to calculate C_{wax} as described in supplementary information **a**. C_{wax} of the two oils are shown in **Figure 2**.

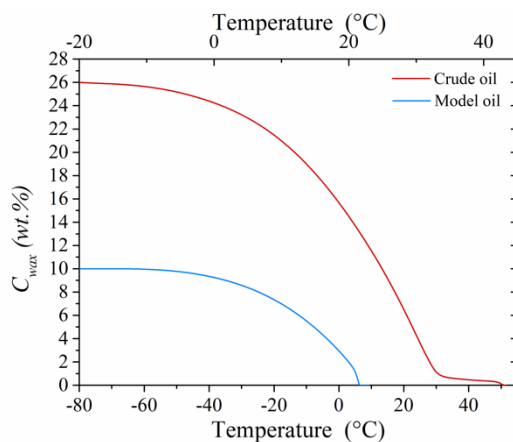


Figure 2. Evolution of the mass fraction of wax crystals C_{wax} in the crude oil from 45°C to – 80 °C (bottom X axis) and the model oil (top X axis) from 20°C to – 20 °C.

2.2. Liquid separation from the solid/liquid gelled mixture by temperature-controlled centrifugation

To better understand the liquid phase behavior during the crystallization process, the liquid phase was extracted at various temperatures from the solid/liquid gelled mixture using a temperature-controlled centrifugation method at five different temperatures. Indeed, this method allows to separate the liquid phase from the *n*-paraffins crystals and to study its composition. The experimental protocol is described in supplementary information e. After the experiments, five centrifuged crude oils with different WAT were obtained (**Table 3**). Their compositions are given in supplementary information c.

Table 3. Centrifuged oils samples obtained at different temperatures with their associated WAT.

	Oil 40	Oil 20	Oil 5	Oil 0	Oil -5
WAT (°C)	32.7	15.1	1.3	-3.4	-9.7

2.3. Cross Polarized Microscopy (CPM)

CPM was used to visualize directly the size and the shape of wax crystals. CPM imaging was performed with an Olympus BH2 UMA microscope, which was fitted with cross-polarization filters and an AxioCam 305 color camera. Temperature and cooling rate were controlled by a LNP95 Linkam. The samples were preheated, filled into glass capillaries with a cross section of 0.2 mm x 4 mm and were sealed on both ends using glue. We recorded two images, using different objectives ($\times 10$, $\times 20$), to compare the wax crystals sizes obtained respectively for the crude oil and for the model oil. During measurements, the temperature was first ramped up to 70 °C (crude oil) or 40 °C (model oil), kept constant for 5 minutes and then decreased down to 0 °C (crude oil) or 18 °C, 5 °C and -10 °C (model oil) at a rate of -10 °C/min. The temperature was held constant at 0 °C or 18 °C, 5 °C and 10 °C and images were taken during the isothermal period.

2.4. Low field Nuclear Magnetic Resonance (NMR)^{39,40}

During the crystallization of *n*-paraffins, the network of crystals formed in the liquid medium can be shown as a porous network (also referred as a crystals network). This porous network consists of distinct pores which contains the liquid that is more or less confined during crystallization. NMR is a non-destructive technique that allowed us to get indirect information about the pore sizes and the behavior of the liquid within the pores using the T_2 relaxation time distribution.

NMR experiments were conducted on a low field (MQR GeoSpec core analyzers: 20 MHz) Oxford Instrument spectrometer equipped with a 18 mm diameter probe. To acquire the transverse relaxation time (T_2) decay, a modified Carl-Purcell-Meiboom-Gill (CPMG) sequence was used: the inter echo spacing (2τ) varies from 100 μ s to 300 μ s and the number of echoes acquired was set to 20 000. The T_2 relaxation time distributions were calculated using the IFPEN

software MEA (Multi-Exponential-Analysis). Glass tubes suitable for the 18 mm NMR probe are filled with approximately 1 g of the fluid sample and then put in the NMR probe. Nitrogen was used to heat and cool the sample. To control the temperature of the sample, the spectrometer was connected to a VT (Variable Temperature) controller coupled with a FTS Low Temperature Cooling Systems & Equipment that allow us to performed measurements over a temperature range between 80 °C and -20 °C. The temperature is maintained by the circulation of a flow of air. To ensure that the sample temperature was stable, the evolution of the magnetization $M(t)$, as a function of time, was followed after several Hahn Echoes (a Hahn Echo is a CPMG sequence with one echo). The temperature of the sample was considered stable when the magnetization remains constant. The stabilization time was about 30-45 minutes. Each raw curve obtained was normalized by the mass of the sample in the glass tube.

In a homogenous B_0 field, we followed the magnetization decay $M(t)$ in the plane transverse to the B_0 field, which is characterized by the relaxation time T_2 . This characteristic time results essentially from liquid-solid interactions of the nuclear spins carried by molecules exploring the pore space by diffusion. Molecular diffusion leads to an exchange between the surface and bulk volumes.

Assuming uniform surface relaxivity and a rapid exchange between liquid and all crystals surfaces, the relaxation time T_2 can be expressed as⁴⁰:

$$\frac{1}{T_2} = \frac{1}{T_{2B}} + \rho_2 \left[\frac{S}{V} \right]_{NMR} \quad (1)$$

Where T_{2B} is the transversal relaxation time of the bulk liquid, V and S are respectively the volume and the surface of the total pore space. ρ_2 is the surface relaxivity or the relaxation velocity at the pore surface. For a uniform pore network, the volume to surface $[V/S]_{NMR}$ ratio gives an estimation of the pore size.

In the presence of a distribution of pore sizes, and in the regime of fast molecular exchange bulk-surface in each pore but not between pores, the measured magnetization decay $M(t)$ is analyzed in term of a discrete series of exponential decays such as:

$$M(t) = \sum_i A_i \exp\left(-\frac{t}{T_{2i}}\right) \quad (2)$$

Where A_i is the normalized number of spins associated with a relaxation time T_{2i} . The T_2 relaxation time distributions are obtained by plotting a curve $A(T_2)$. The total magnetization $M(t=0)$ represents the total amount of spins. In practice, it is possible to detect only the spins that have a sufficiently long relaxation time: this is the case of the bulk liquid. On the contrary, for solids, relaxation times are very short and the detection of the magnetization associated with these spins will depend on the spectrometer and its characteristics. For example, during our CPMG measurements performed at 20 MHz we cannot detect the magnetization associated with n -paraffins crystals, which have a relaxation time T_2 of 70 μs ²¹. $M(t=0)$ varies according to the inverse of temperature. To correct the effects related to temperature variation, the measured magnetization is corrected by the Curie's law.

2.5. Small and Wide Angle X-ray scattering (SAXS/WAXS)

SAXS/WAXS were used to investigate the crystallographic structure of n -paraffins crystals, their form, size and number density, their surface area as well as their spatial correlation at larger length scale. SAXS/WAXS experiments were performed at IFPEN. The experimental set-up is equipped with a copper rotating anode (Rigaku MM07) providing a X-ray beam with a wavelength of $\lambda = 1.5418 \text{ \AA}$. When the beam crosses the sample, a small part is scattered and is captured simultaneously by a SAXS and WAXS two-dimensional (2D) detectors (respectively Rigaku and Xenocs). This design allows to collect scattering intensities over a large range of

scattering angles 2θ . Each pixel coordinate was converted to wave vector q thanks to measurement of standards, namely Ag Behenate and LaB₆ respectively for SAXS and WAXS⁴¹. The scattering vector q is defined as $q = \frac{4\pi\sin(\theta)}{\lambda}$, with θ being half the deviation angle, and corresponds to an inverse correlation length probed. In the case of a periodic structure, a value of q in the reciprocal space is related to a repeating distance d in the real space by the Bragg law: $d = 2\pi/q$. For experiments, working with two different distances between sample and detectors allows to cover a total q domain of almost three decades ranging from $6 \cdot 10^{-3} \text{ \AA}^{-1}$ up to $4.3 \text{ \AA}^{-1}$. SAXS and WAXS experiments were carried out simultaneously on the same sample volume. The crude oil and the model oil were preheated and filled into glass capillaries with a diameter of 1.5 mm, which were flame-sealed on both ends. Capillaries were then placed on the Linkam temperature-controlled sample holder and heated for a few minutes to 70 °C (crude oil) or 50 °C (model oil) before the measurements begin. The whole stage was placed in a vacuum chamber to avoid any condensation or frost on wall capillaries upon cooling. Then, the temperature was cooled down at 10 °C/min from 70 °C to -70 °C or from 50 °C down to -20 °C. The raw scattered intensities were normalized by the thickness of the sample, the transmission and the measuring time. Empty capillary as well as liquid intensities, obtained from Hydroseal or centrifuged oils above their WAT, were subtracted to each measurement according to their volume fraction. Finally, Lupolen was chosen as calibration standard⁴² to convert intensities in absolute units (cm^{-1}).

For a two-phase system made of monodisperse particles distributed in space, the general equation describing the scattered intensity of particles in the solvent can be written as:

$$I(q) = N_p \Delta\rho_{scat}^2 v^2 P(q) S(q) \quad (3)$$

Where N_p is the number density of scattering particles and v is the volume of one particle. N_p can be written as ϕ/v where ϕ is the particle volume fraction. $P(q)$ is the form factor, which is a function of shape and size of particle [$P(0)=1$]. $S(q)$ is the structure factor describing particles position correlation resulting from their mutual interactions. For concentrated systems, the structure factor provides significative structural information at length scale larger than the typical size R_G of scatterers, i.e at q values smaller than $\sim R_G^{-1}$. At q values much larger than R_G^{-1} , $S(q)$ tends to 1. $\Delta\rho_{scat}$ is a contrast term, which is the scattering length density difference between the particles and solvent.

The form factor $P(q)$ can be expressed in different ways depending on the particles shapes and the q domain:

At large q values ($q > 1/l$), with l being the smallest dimension of the particle, known as the ‘Porod’ regime, it is possible to determine the total surface area developed by particles (S) in a given volume (V) using the following equation:

$$I(q \rightarrow \infty) = \Delta\rho^2 \frac{2\pi}{q^4} \left[\frac{S}{V} \right]_{SAXS} \quad (4)$$

The surface to volume ratio $[S/V]_{SAXS}$ can also be written as:

$$\left[\frac{S}{V} \right]_{SAXS} = \frac{s}{v} \times \phi \quad (5)$$

Where s and v stand respectively for the surface area and the volume of one particle and ϕ is the particle volume fraction.

At intermediate q values, the variation of the scattering intensity $I(q)$ can follow a power law defined as:

$$I(q) \sim q^{-\alpha} \quad (6)$$

Where α is related to the dimensionality of the structure. In the dilute regime, $\alpha = 1$ for rods and 2 for lamellae or discs.

The size of crystalline domains S_{hkl} , in a direction perpendicular to a $h k l$ atomic plane, represent the size of ‘perfect’ crystal (without crystalline defects), they give a minimal value of the particle size in the considered direction. They are inferred from the full width at half maximum $FW(S)_{hkl}$ of the considered (hkl) Bragg reflection through the Scherrer relation described in supplementary information **g**.

3. RESULTS

3.1. Crystals observations

To get a first estimation of the *n*-paraffins particles form and size at the micrometric scale, CPM experiments were conducted. At high temperatures, only a homogeneous black background, corresponding to the solvent, is visible. When the sample is cooled and temperature falls below the WAT, bright white particles appear as a result of birefringent crystal formation. For the crude oil, the perceptible particles are small (size of a bright spot $\approx 1 \mu\text{m}$) and it is difficult to have an accurate information about their size and shape (**Figure 3a**). On the contrary, for the model oil, it is easier to distinguish the different *n*-paraffins crystals and to determine their size and shape (**Figure 3b** and **Figure 3c**). Crystals appear in various orientations (**Figure 3b**), some of them being positioned flat or on the slice. They are thin platelets made up of a stack of lamellae with typical dimensions (150 μm length x 50 μm width x 5 μm thickness). The areas pointed by arrows in **Figure 3b** and **Figure 3c** shows the stacking of lamellae constituting the platelets. These lamellae seem to divide into several branches and form a less concentrated area of

275 lamellae (zoom **Figure 3b**). As the temperature decreases, the lamellae divide more and more
 276 into several branches and form a structure called hedrite (**Figure 3c** and **Figure 3d**). The hedrite-
 277 shaped structure is a structure observed during the crystallization of semi-crystalline polymers⁴³.

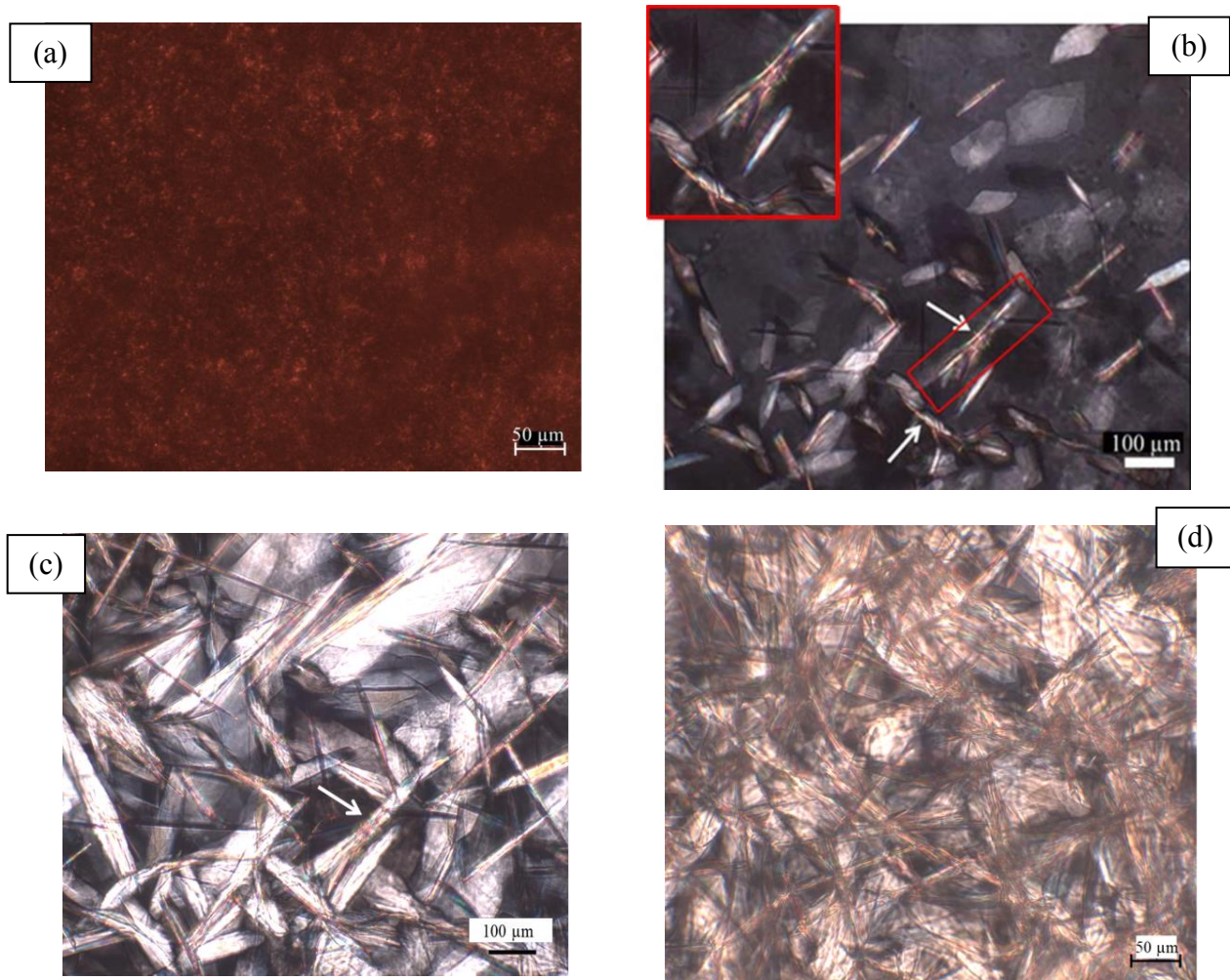


Figure 3. CPM images obtained a) at 0°C ($\times 20$ magnification) for the crude oil (WAT = 51°C). For the model oil (WAT = 22°C) b) at 18 °C ($\times 10$ magnification) c) at 5 °C ($\times 10$ magnification) and d) at -10 °C ($\times 20$ magnification) after cooling down at -10 °C/min. The areas pointed by arrows represent the aggregate which is composed of a stack of crystals that divide into branches.

3.2. From *n*-paraffins chains to crystals aggregates

To go further in the characterization of crystals, we now analyze SAXS/WAXS results. The 2D scattering patterns (images are not shown) obtained appears fully isotropic for the crude oil and slightly punctuated for the model oil. The punctuation of Bragg rings appears when all the crystals orientation in the probed volume are not equally probable. From CPM observations, we suspect that crystals in the model oil grew to such large size that some preferential orientation occurs. However, as the rings were quite homogeneous and almost isotropic, images were azimuthally averaged to obtain 1D curves.

In order to represent only the major evolutions and to allow a good readability of the results, we represent in **Figure 4** the recomposed 1D spectra, in the different q domains, obtained at only a few temperatures above and below the WAT. They are analyzed according to (i) their Bragg peak features ascribed to phase identification and crystal size domains and to (ii) their general behavior at small q values.

3.2.1. Crystalline structure

At wide angles, and above the WAT, the X-ray scattering spectra in **Figure 4** present a wide halo located at $q = 1.31 \text{ \AA}^{-1} \pm 0.01 \text{ \AA}^{-1}$ ($d = 4.5 \pm 0.03 \text{ \AA}$). This halo is associated with the liquid phase. As expected, the halo area decreases below the WAT. When the temperature falls below the WAT, four narrow Bragg peaks appear, located at $d = 4.1 \text{ \AA} \pm 0.03 \text{ \AA}$, $d = 3.7 \text{ \AA} \pm 0.02 \text{ \AA}$, $d = 3.1 \text{ \AA} \pm 0.02 \text{ \AA}$ and $d = 2.4 \text{ \AA} \pm 0.01 \text{ \AA}$. These peaks are respectively attributed to the $hk0$ reflections, (110+111), (020), (120) and (200), of an orthorhombic crystal system^{44,45}. The most intense one appears just at the WAT temperature, which confirms the sensitivity of X-ray scattering for the Wax Appearance Temperature determination³⁰.

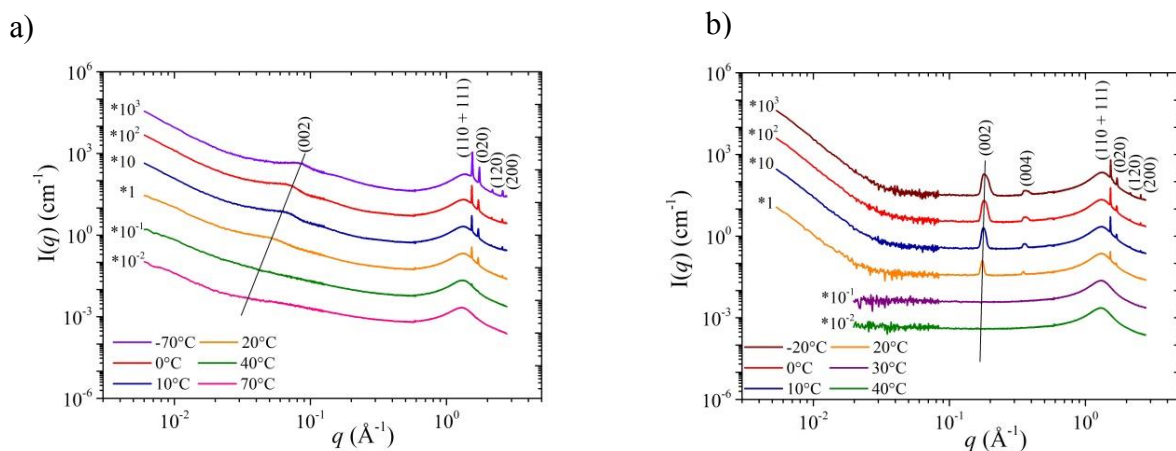


Figure 4. X-ray scattering spectra at different temperatures before the solvent subtraction for a) the crude oil (WAT = 51 °C) and b) the model oil (WAT = 22 °C).

The unit cell parameters a and b are inferred from peak positions and indexations using equation described in supplementary information **g**. For the two oils, these parameters slightly decrease with temperature (**Figure 5**). This small decrease is of the same order of magnitude than the one reported for pure alkanes samples⁴⁶, which remain in an ordered orthorhombic crystalline phase with no phase transition. There is no other Bragg peak appearance in the q domain under study, excluding the possibility of another low temperature phase transition.

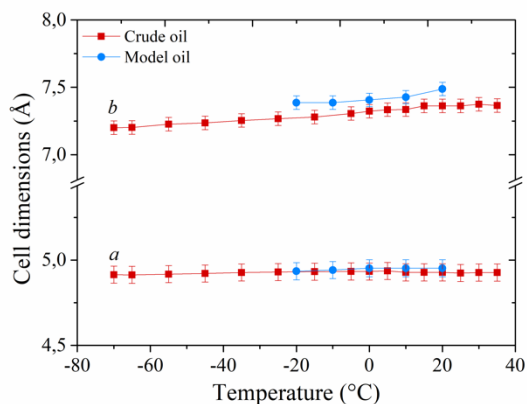


Figure 5. Temperature dependence of the lateral unit cell parameters a and b for the crude oil and the model oil.

At small angles, and below the WAT, the X-ray scattering spectra in **Figure 4** show one peak for the crude oil and two (harmonic) peaks for the model oil located at different positions depending on the oil and temperature. They are indexed as the (002) – and (004) – Bragg peaks of the orthorhombic unit cell^{44,45} and are related to the spacing between perpendicular planes to the chains axes (or the c direction). Their positions are related to the mean length of molecular chains incorporated in crystals. Indeed, the simple relation $c (\text{\AA}) = 2 d_{002} = 2.544 \overline{n_{RX}} + 3.75$ has been proposed, where $\overline{n_{RX}}$ stands for the mean carbon atom number chain incorporated into crystals^{35,38,47}. For the crude oil, at 20 °C, $\overline{n_{RX}}$ is 50, while for the model oil, at the same temperature, $\overline{n_{RX}}$ is 27. These results highlights the differences in the n -paraffins distribution of the two oils.

The previous relation is valid for several situations²⁶ including the pure alkane phases up to solid solutions for n -alkane mixture of a low carbon atom number polydispersity σ of 1.5. For larger polydispersity ($3 < \sigma < 4.4$), $\overline{n_{RX}}$ is one or two unity larger than the prevalent carbon number in the mixture. For even more polydisperse synthetic waxes ($\sigma = 5.8$), three different solid solutions are identified at room temperature and disappear upon melting. The high temperature solid solution crystallizes chains with mean carbon atom number $\overline{n_{RX}}$ much larger than the prevalent carbon atom number $\bar{n}$ of synthetic wax ($\overline{n_{RX}} - \bar{n} = 7$). At the opposite, the low temperature solid solution crystallize chains shorter than the prevalent one ($\overline{n_{RX}} - \bar{n} = 2$).

By analogy, we could infer, from the polydispersities of the model oil and crude oil, that the former will crystallize in a single solid solution whereas the latter will crystallize in several solid solutions.

3.2.2. Surface to volume $[S/V]_{SAXS}$ ratios

At small q values, the crude oil spectra (**Figure 4a**) show a significant scattering signal at high temperatures (above the WAT) while the model oil spectra (**Figure 4b**) present a flat signal. The flat signal is related to a ‘true’ liquid (*i.e.* an assembly of small molecules) whereas the q dependent signal is associated with electronic density fluctuations at $\sim q^{-1}$ length scale. Such upturn of scattering intensity at small q values has already been noticed on de-asphalted crude oils^{48,49} without any further investigations on the nature of such fluctuations. The crude oil subtracted curves (**Figure 6a**) show a first regime at low q with a q^{-2} dependence and a second one at large q following a q^{-4} power law. The q^{-4} dependence (Porod domain) allows to estimate the surface to volume $[S/V]_{SAXS}$ ratio according to the equation (3). The surface to volume $[S/V]_{SAXS}$ values are plotted in **Figure 7a**. Values are quite high, in the range of $10^5 - 10^6 \text{ cm}^{-1}$, and vary in the same way as the crystal content as a function of temperature. For the model oil, we observe a q^{-4} power law in the whole q range (**Figure 6b**). As for the crude oil, this *Porod* regime allows to estimate the surface to volume $[S/V]_{SAXS}$ ratios (**Figure 7b**). The surface to volume $[S/V]_{SAXS}$ ratios vary in the same way as the crystal content but their values, $0.8 \text{ to } 2 \cdot 10^3 \text{ cm}^{-1}$, are three decades lower than for the crude oil.

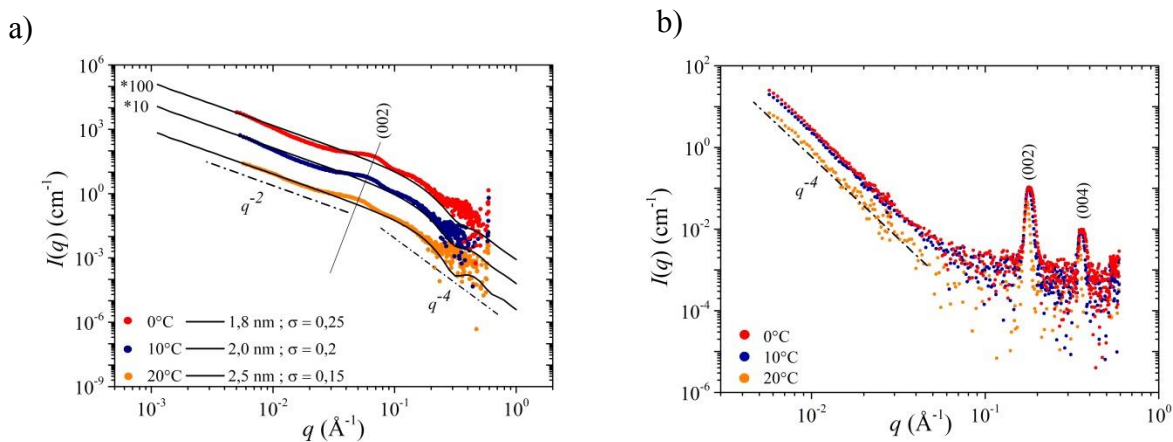


Figure 6 . a) crude oil and b) model oil SAXS spectra from Figure 4 subtracted from solvent and normalized by the sample volume fraction. The solid lines in Figure 6a correspond to the fit of a flat disc model of variable polydisperse thickness.

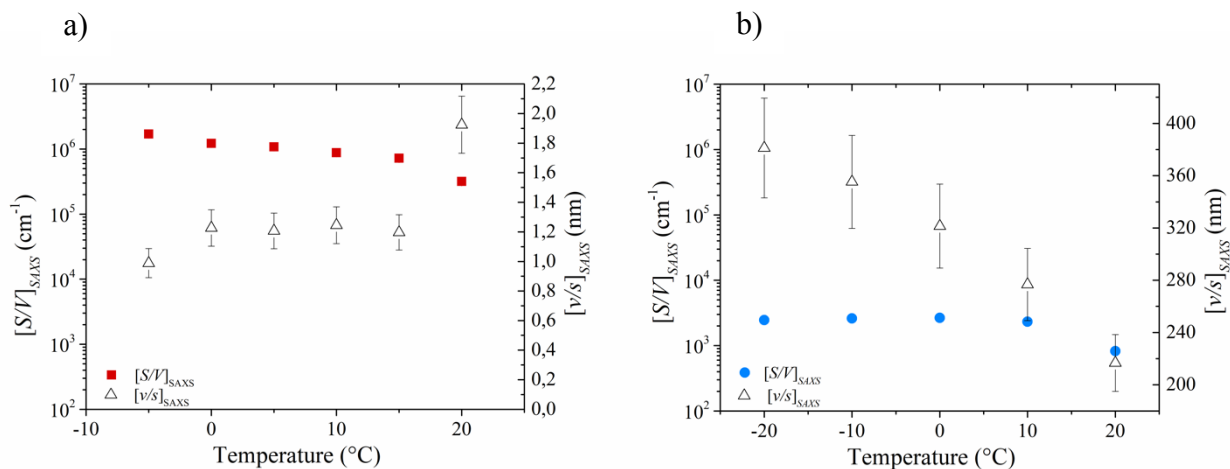


Figure 7. Surface to volume $[S/V]_{\text{SAXS}}$ and volume to surface $[v/s]_{\text{SAXS}}$ ratios obtained from SAXS measurements for a) the crude oil and b) the model oil.

3.2.3. Crystals form

The q^{-2} behavior observed on the crude oil spectra (**Figure 6a**) is reminiscent of 2D objects such as discs or lamellae.

3.2.4. Crystals size

The crystal size domain in direction a or b perpendicular to chain axis, is related to the width of the $(hk0)$ peak at half height. For the two oils, the measured widths of the (110) peak are in the same order of magnitude as the instrument resolution (**Figure 8**). It means that the crystal size domains, in the crude oil and the model oil, are larger than $\approx 1000 \text{ \AA}$ in direction a and b . So, in the directions perpendicular to chain axis, crystals show a long-range order.

Then, $(00l)$ reflections can be used to measure the crystals size domains in the direction of the chains on the whole range of temperature for the model oil and just below the WAT for the crude oil. For the two oils, the measured widths of the (002) peak do not have the same order of magnitude as the instrument resolution (**Figure 8**). The maximum crystallite size, in direction c , is found to be of the order of $160 \text{ \AA}$ in the crude oil and $1200 \text{ \AA}$ in the model oil. These sizes are typical of anisotropic objects, which is in good agreement with lamellar crystals.

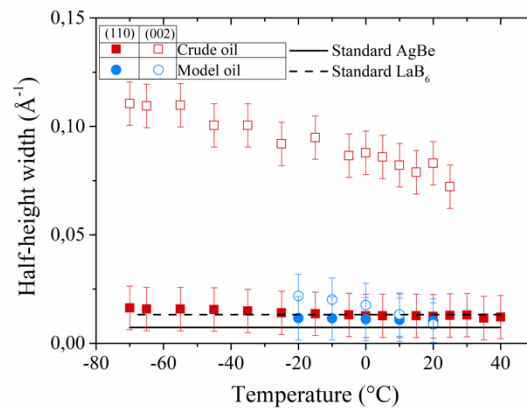


Figure 8. Half-height width of the peaks (110) and (002) of the crude oil and model oil as a function of temperature.

For flat particles of thickness $2H$, the volume to surface ratio $[v/s]$ of one particle is simply $2H$, and could be obtained from the total surface to volume $[S/V]_{\text{SAXS}}$ ratio and the volume fraction ϕ (equation (5)). For the crude oil, $2H=v/s$ are plotted as a function of temperature on **Figure 7a**.

The $2H$ values are quite small, in the range of monomolecular thickness, and seems rather constant in the studied temperature range, suggesting that each new crystal formed has a monomolecular thickness^{20,28,50}. To confirm this thickness and since the interactions between particles are not probed in the observed q range, the whole SAXS patterns are fitted to flat discs form factor (equations are described in supplementary information f). The results (**Figure 6a**) indicate a mean thickness of 2-3 nm, which is in good agreement with the thickness deduced from the surface to volume $[S/V]_{\text{SAXS}}$ ratios. So, in the direction of chain axes, crystals from the crude oil develop a very short range order.

The cross over point, q^* , defined as $q^* = 1/2H$, between the q^{-2} and q^{-4} behavior can also be used to estimate directly the thickness $2H$ of the particles. From the *Porod* regime observed in the model oil spectra (**Figure 6b**), it is possible to obtain a minimum value of $2H_{\min}$ ($2H_{\min} > 1/q_{\min}$) for the model oil. The minimum value of $2H_{\min}$ found is approximately 160 Å.

If we assume, as suggested by CPM and Bragg peak broadening observations, that model oil particles are lamellae, we can also estimate the crystal thickness $2H=v/s$ (**Figure 7b**). Their values increases slightly as temperature decrease but we can retain an average value of $2H_{(v/s)} \approx 3000$ Å. This thickness is (i) larger than the crystal size domain (1200 Å) highlighting likely the stacking defect in the chain axis direction, (ii) smaller than the CPM observation that is also questionable in respect to parallax effects and to resolution, (iii) greater than the minimum value of $2H_{\min}$ found. Anyway, the lamella thickness for the model oil is 2-3 order of magnitude larger than for the crude oil showing a longer-range order.

3.3. Fluid dynamics within the crystals network

Information about the fluid dynamics, its state of confinement within the crystal network formed are extracted from NMR results.

391 The T_2 distribution curves obtained at few temperatures, from above the WAT detected by DSC
392 (51 °C and 22°C) to below the WAT, for the two oils (after the multi-component decomposition)
393 are shown on **Figure 9**. Concerning the crude oil, as the amount of crystals is low between 51 °C
394 and 31 °C, NMR detects the appearance of crystals and a change in behavior at a temperature
395 $T_{\text{crystals}} = 30$ °C. For the model oil, the change in behavior and the appearance of the crystals
396 coincide with the WAT detected by DSC ($T_{\text{crystals}} = 22$ °C).

397 Above T_{crystals} , the T_2 distributions show a main peak with high T_2 values (**Figure 9**). For the
398 crude oil, the peak is wide and asymmetrical, while for model oil it is thin and symmetrical. The
399 asymmetrical distribution shape of the peak has already been observed on de-asphalted crude
400 oil⁵¹ and is probably due to the presence of objects in the liquid phase which are remaining above
401 the WAT, as observed on the crude oil SAXS spectra (section 3.2.2).

402 As the temperature decreases, the two oils show similar behaviors since the main peak shifts to
403 shorter T_2 and expands. The shift and expansion are caused by an increase of the bulk oil
404 viscosity with the temperature and by the formation of surfaces magnetic relaxations that appear
405 during crystallization.

406 Below T_{crystals} , the crude oil results show the presence of a visible shoulder and peaks at short T_2
407 that does not exist in the model oil. The shoulder intensity increases and the intensity of the main
408 peak decreases as the temperature decreases. These behaviors can be related to SAXS results and
409 reflect a high quantity of surfaces formed (**Figure 7**, section 3.2.2) and a reduction in the bulk oil
410 protons mobility, which is more significant in the crude oil compared with the model oil. From
411 these observations, we can assume that there are two different protons populations with different
412 mobility, which interact with the surfaces created during crystals formation. They interact with

413 two different environments and with different characteristics, including different solvent domain
 414 sizes.

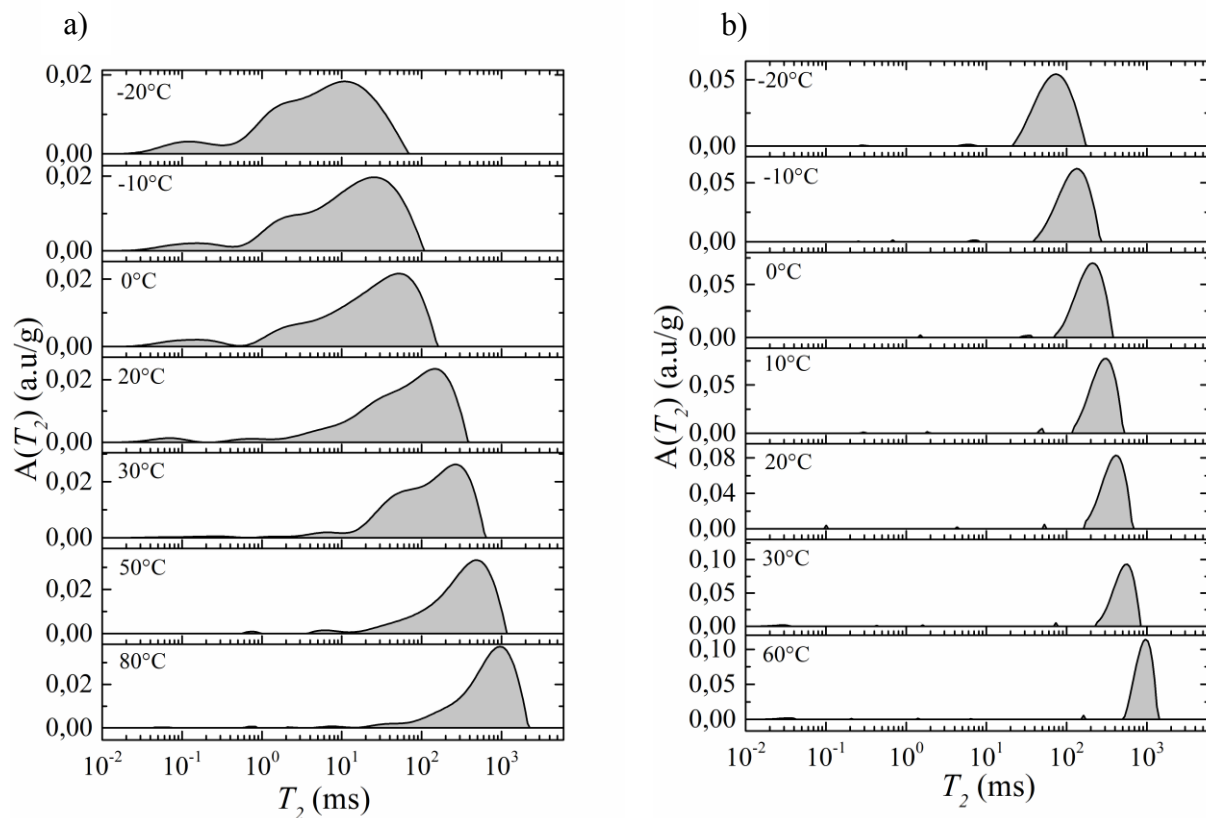


Figure 9. Relaxation time distributions, plotted as the log relaxation time versus the normalized signal intensity, at different temperatures for a) the crude oil and b) the model oil. The relaxation time distributions shown were obtained at an echo time of 100 μ s.

415 3.3.1. NMR sensitivity to crystals surfaces

416 We are now interested in the crude oil and model oil main peak evolution present at large T_2 as
 417 the temperature decreases (**Figure 10**). We considered the maximum of the distribution as a
 418 tracer of T_2 evolution.

419 Above T_{crystals} , the function $\ln(T_2)$ varies linearly with the inverse of the temperature. A straight
 420 line is drawn to show this evolution. When the temperature decreases, T_2 decreases until T_{crystals}

and then deviates from this first line. This deviation is due to the creation of crystals and new surfaces relaxation. To confirm that, measurements on oil 20 and oil 5 were conducted above their WAT. The results show that the T_2 obtained are in line with the T_2 of the crude oil obtained at $T > T_{\text{crystals}}$ and confirm that the deviation is due to the formation of surfaces in the mixture (Figure 10).

Protons of the liquid phase are now interacting with the spins of surfaces. It can be noticed that the deviation is more important for the crude oil than for the model oil, due to the large amount of surfaces areas present in the crude oil as observed with SAXS/WAXS results (Figure 7, section 3.2.2).

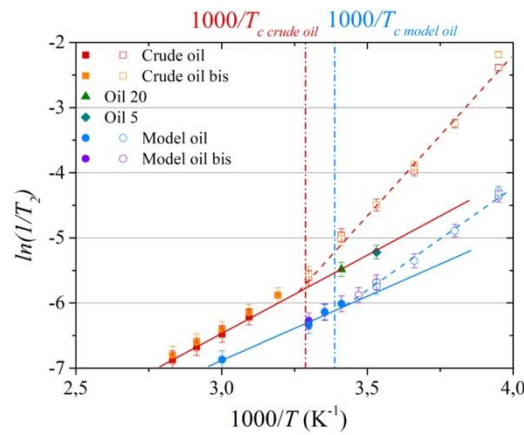


Figure 10. Evolution of the main peak relaxation rate obtained, from the multi-components distributions, versus the inverse of temperature for the crude oil and the model oil. “Bis” refers to a repetition of a set of measurements.

3.3.2. Confined and unconfined liquid

Then, to go further in understanding the dynamic process and proton populations associated with this system, we focused only on the main peak and its shoulder by decomposing the T_2 distributions into only two components. The first three points of the distributions were removed in order to only take into account the contribution of the liquid which relaxes at $T_2 > 0.1$ ms

435 (supplementary information **e**). The two-components decomposition is carried out at all
 436 temperatures below T_{crystals} . An example of the decomposition and a fit with a lognormal
 437 distribution, on the signal at 0 °C, is represented on **Figure 11**. The main peak is represented by
 438 the decomposition (1) and the shoulder by the decomposition (2).

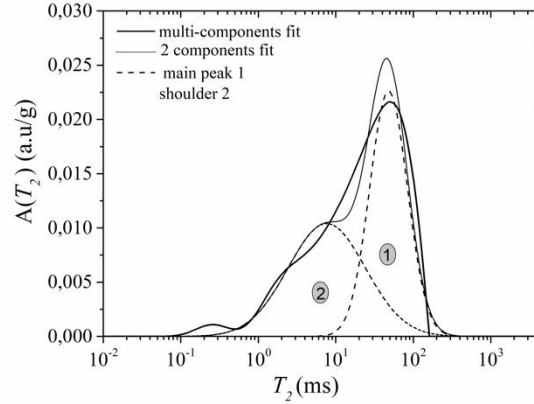


Figure 11. The crude oil relaxation time distribution obtained at 0°C with a multi-component fit (dark solid line) and a two-component fit (light solid line). The two-component distribution (light solid line) is decomposed into two peaks (dashed line): the main peak represented by component (1) and the shoulder represented by component (2).

439 The peaks proportion evolution was studied to characterize the exchanges that can take place
 440 between the two peaks and have more information on the behavior of each proton population as
 441 the temperature decreases (**Figure 12**). The two-components fit gives the proportion of the main
 442 peak and the shoulder associated with the measured magnetization M_{measured} . The total proportion
 443 of the main peak and shoulder (1+2) corresponds to the ratio between the magnetization
 444 measured after decomposition into two components and the magnetization extrapolated by the
 445 Curie's law $[M_{\text{measured}}/M_{\text{extrapolated}}]$.

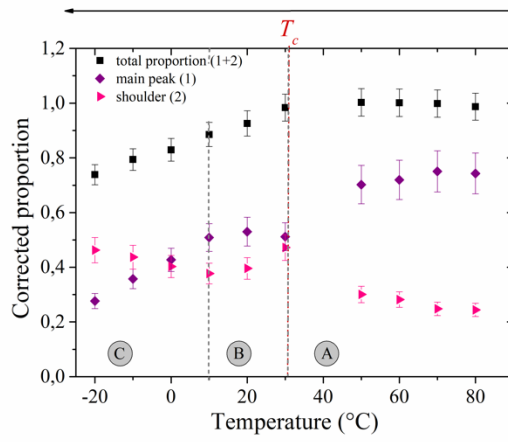


Figure 12. Corrected proportions of extrapolated magnetization attributed to the main peak and shoulder as a function of temperature (see text for detailed explanation). A,B,C represent the different behavior observed. These behaviors are explained in the text.

On **Figure 12**, we can notice that the shoulder is pre-existing at high temperatures and its proportion is not negligible ($\approx 20\%$). Three regimes can be distinguished when the temperature decreases:

- In the first regime A ($T > T_{\text{crystals}}$), the total proportion remains constant and we observe an increases of the shoulder proportion (2) while the proportion of the main peak (1) decreases. This can be related to the n -paraffins molecules that are less and less mobile.

- In the transition regime B ($T < T_{\text{crystals}}$), the main peak proportion (1) stays stable, the shoulder proportion (2) slightly decreases and the total decreases. This is due to the creation of n -paraffins crystals. Indeed, a part of protons initially associated to the shoulder are now incorporated in solids.

- In the final regime C ($T \ll T_{\text{crystals}}$), the total and main peak proportion (1) decreases whereas the shoulder proportion (2) increases. There is an important part of the main peak proton population (1) which is now associated with the shoulder protons environment. This means that in addition to the creation of crystals, there is also a lot of new surfaces created. This observation is consistent with $[S/V]_{\text{SAXS}}$ SAXS results.

These observations confirm that between the two population of protons identified by NMR, the main peak corresponds to a liquid less confined than the one represented by the shoulder. Therefore, to go further, one would like to evaluate the liquid distances between the surfaces associated with these two proton populations identified. From the surface to volume $[S/V]_{SAXS}$ ratio values obtained a mean liquid distance over the entire sample d_{mean} can be obtained from equation (7) :

$$d_{mean} = \frac{1}{\left[\frac{S}{V}\right]_{SAXS} \left(\frac{1}{(1-\phi)}\right)} \quad (7)$$

The values of d_{mean} obtained (**Figure 13**) are on the order of few nanometers. However, as an order of magnitude, the chain length of an *n*-paraffin molecule containing 15 carbons is 4 nm, which is of the same order of magnitude as d_{mean} . This means that the liquid is very confined with few or no bulk volume and there is mostly adsorption of the molecules on surfaces. In this case, the fast exchange model is not valid. Thus, the distances associated with each protons population cannot be determined from the equation (1) and surface to volume $[S/V]$ ratios obtained by NMR and SAXS as presented in supplementary information f.

Ongoing studies could thus focus on further information about the two protons population identified. On the one hand, at large scale, the exchange dynamics between the different proton populations could be investigated by (2D) NMR T_2 - Δ - T_2 ⁵². On the other hand, it would be interesting to probe by NMRD⁵³ the local interfacial dynamics of these protons populations in these pronounced confinements.

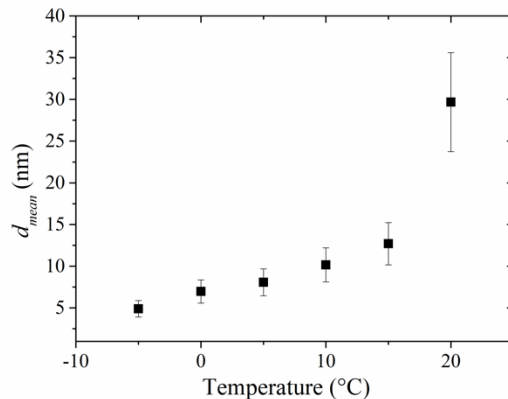


Figure 13. Liquid distances between surfaces d_{mean} estimated from SAXS measurements as a function of temperature.

4. DISCUSSIONS AND CONCLUSIONS

In the current study, we focused on n -paraffins crystallization to probe at multi length-scale the n -paraffins structure and the behavior of the liquid within the crystal network formed during crystallization. A multi modal strategy was proposed including CPM, SAXS/WAXS and low field NMR experiments. Two waxy oils, a crude oil and a model oil, of different n -paraffins polydispersities and compositions were used. The major results of the study are summarized in **Table 4** below.

Table 4. Summary of the major results ($n.d$ = not determined).

	n -paraffins polydispersity σ	Crystalline structure	Crystals thickness (nm)	Crystals shape	$[S/V]_{SAXS}$ (cm^{-1})	Liquid distance d_{mean} (nm)
Crude oil	>7.2	Orthorhombic	1-10	Lamellar	$\approx 10^6$	5-30
Model oil	2.1	Orthorhombic	> 120	Lamellar	$\approx 10^3$	n.d

Above the WAT, no structure should subsist since the n -paraffins are totally solubilized. Though, no singular behavior is observed for the model oil. The results obtained with the crude oil by SAXS and NMR revealed the presence of molecules at high temperatures with a distinct signature, which were unobservable by CPM. Asphaltenes but also other complex molecules could be at the origin of such behavior as reported in the literature on de-asphalted crude oils^{48,51}.

495 More investigations, out of the scope of the present paper, could be performed to probe which
496 molecules are responsible for this peculiar signature.

497 Below the WAT, *n*-paraffins crystals appear. In the model oil, crystals form a single solid
498 solution with an orthorhombic structure. In the crude oil, several solid solutions with an
499 orthorhombic structure are formed. In both cases, the cell parameters are about $a = 7.40 \text{ \AA} \pm 0.05$
500 $\text{\AA}$ and $b = 4.92 \text{ \AA} \pm 0.05 \text{ \AA}$. The *n*-paraffins crystals size is about 1-10 nm in the crude oil and
501 larger than 120 nm in the model oil. These differences in size could be due to the presence of
502 non-crystallizable molecules such as isoalkanes or cycloalkanes, resins and asphaltenes, which
503 inhibit crystal development and promote the formation of small structures⁵⁴⁻⁵⁶. A high
504 concentration of resin-type molecules can lead to aggregates of poorly crystallized forms that
505 grow following a branch-like shape. As crystals grow, they change direction because the
506 localized accumulation of impurities on surfaces promote the formation of branches⁵⁶. Studies on
507 long *n*-paraffin chains and semi-crystalline polymers have also shown that the structure of crystal
508 aggregates depends on the applied undercooling ($\Delta T = T - T_c$ with T_c the crystallization
509 temperature)⁵⁷. At low degree of undercooling the crystal aggregates will be more hedritic in
510 shape and at larger degree of undercooling the branches will divide more and more and the shape
511 of the aggregate will tend more towards spherulites, which are a set of thin lamellae organized in
512 spherical and symmetrical macrostructures⁴³. These conclusions drawn from the literature can be
513 compared with the results we obtained in this study. Indeed, SAXS measurements showed that
514 the formed crystals have a lamellar shape. Thus, we can deduce from the model oil CPM images
515 that the observed platelets composed of a stack of lamellae are in fact aggregates of crystals of
516 lamellar shape. As observed in the literature^{43,56}, we could expect that they grow in branched
517 form with branches that divide more and more as the temperature decreases. However, since the

model oil does not contain asphaltenes and resins, the crystals are larger and the branches divide significantly less than in the crude oil (**Figure 14**). This observation can be supported looking at the evolution of trapped liquid protons population identified in the crude oil and unobservable in the model oil: the more branches, the greater the amount of liquid trapped between the branches. Future NMR studies could give more information about the dynamics of these trapping liquid areas. Furthermore, it is worth to notice that during the formation of the branched *n*-paraffins crystals, liquid is trapped and the effective volume of the system (crude oil + crystals) occupies a larger volume than the net volume occupied by the crystals packed together. This effective volume fraction, which is considered when modeling the gelled waxy crude oil as a suspension of fractal aggregates, would be an interesting data to refine or discriminate the existing rheological models^{6,7,58}.

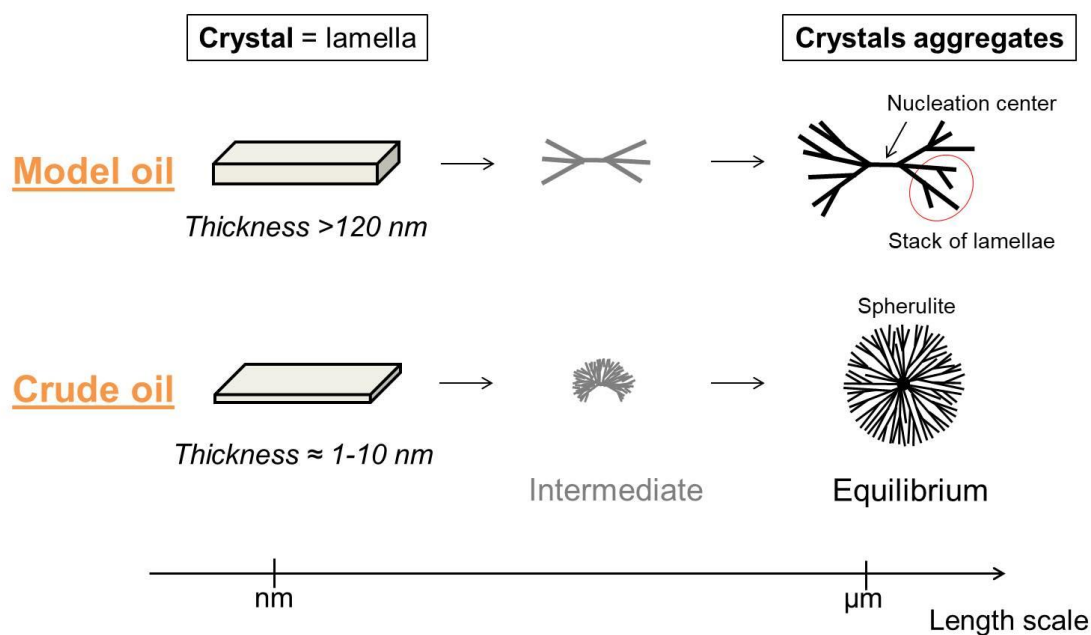


Figure 14. Illustration of the aggregation process of *n*-paraffins crystals in (a) model oil and (b) crude oil.

533

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542 6. SUPPORTING INFORMATION

543
544 The procedure followed to determine the WAT and the WPC is given in supplementary
545 information as well as densities and elemental composition of the oils. Equations used to
546 calculate σ and $\bar{n}$ parameters can be found in supplementary data. The experimental protocol of
547 the temperature-controlled centrifugation is also given. SAXS/WAXS theoretical background are
548 given in supplementary information. For the crude oil, the magnetization evolution measured by
549 NMR and the methodology followed to calculate the distances between the confined liquid and
550 the surfaces created during crystallization are also detailed.

551

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