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# Fast and accurate selection of surfactants for Enhanced Oil Recovery by Dynamic Salinity-Phase-Inversion (SPI)

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## Abstract

In conventional chemical Enhanced Oil Recovery (EOR), surfactant mixtures must be optimized to obtain the so-called “optimum formulation” characterized by a three-phase behaviour (WIII) and an ultra-low interfacial tension between crude oil and injected water. To attain this condition, discontinuous salinity scans of Surfactant-Oil-Water (SOW) systems are usually performed in a series of sealed pipettes maintained at the reservoir temperature until complete phase separation. Furthermore, some SOW mixtures do not give a WIII, but instead they form viscous phases which are very detrimental for EOR applications.

The study herein describes a novel method called dynamic Salinity-Phase-Inversion (SPI) to quickly and accurately determine the optimum formulation and to anticipate whether the surfactant mixture will provide, by the conventional static method, a WIII or a viscous phase. This method consists in continuously modifying the salinity of a stirred SOW system at WOR in the vicinity of 1, in order to detect the phase inversion of the emulsion by a sudden drop in conductivity.

The effectiveness of the method is illustrated by seeking the optimum formulation for two crude oils in the presence of a mixture of an ionic EOR surfactant (sulfated propoxylated *n*-alcohol) and a nonionic

co-surfactant (ethoxylated *i*-alcohol), with or without alkali. Continuous salinity scan of the stirred SOW system maintained at constant temperature allows the identification of the optimal salinity within 30 minutes. This method therefore significantly accelerates the optimization of surfactant blends suitable for a given oil reservoir.

**Keywords:** Salinity Phase Inversion, Optimum formulation, EOR, Crude oil, Fast screening, Winsor III microemulsion

## 1. Introduction

Various scenarios try to shape the world energy demand in the next decades, taking into account the population growth, estimated at more than 9 billion people in 2040, and the climate change awareness. In particular, the evolution of the oil demand in the next decades is uncertain.

This global energy context along with crude oil prices volatility encourage oil producers to optimize the exploitation of their existing fields and to improve the efficiency of their extraction processes. In average, two-thirds of the oil remain trapped in a reservoir after primary and secondary (water flooding) recoveries. One technique to improve the recovery factor is the chemical Enhanced Oil Recovery (EOR) in which an aqueous solution of surfactants and polymers is injected inside the reservoir [1–3]. While polymers allow improving the mobility ratio, surfactants maximize oil mobilization and flow by reducing capillary forces [4]. To achieve an efficient desaturation, the interfacial tension between oil and water has to be reduced down to ultra-low values ( $\sim 10^{-3}$  mN.m<sup>-1</sup>), which can be achieved by finely tuning the surfactants blend. The conventional SP (Surfactant Polymer) and ASP (Alkali Surfactant Polymer) EOR processes aim at achieving very low residual oil saturations. Aside these processes that might be too demanding for full implementation on some fields, more easily deployable techniques using surfactants can be applied continuously or intermittently to improve oil production. For instance, well injectivity can be significantly improved by removing accumulated hydrocarbons near the wellbore [5]. This can be achieved by injecting low dosages – tens

of ppm of well-suited surfactants. Also, wellbore clean-up operations can be greatly improved by using well-designed mud cake remover formulations [6].

All these surfactant-based processes require to adjust the affinity of the surfactant blend to the oil to be extracted or to be removed. In all cases, the oil nature and the temperature of the process are fixed. One way to achieve the so-called “optimum formulation” is to change the salinity of the aqueous phase in order to modulate the relative affinity of the surfactant blend for water and for the targeted oil [7]. In some cases, because of the field constraints and of the water sources availability, the salinity is fixed also and the surfactants blend itself must then be adjusted to reach the optimum conditions.

In practice, the best surfactant system for a given reservoir is determined by salinity scans at the reservoir temperature and the optimum formulation is visually determined at the salinity for which a balanced Winsor III (noted WIII) microemulsion equilibrated system - containing equal volumes of oil and water - is obtained [8–10]. When the optimal salinity is out of the process-required range or if the Surfactant/Oil/Water (SOW) system does not form WIII systems, the surfactants blend is adjusted and new salinity scans are performed in order to check the phase behaviour over the salinity range. These scans require, in some cases, lengthy equilibrium times to unambiguously identify the phase behaviour with crude oil, which is a restricting factor to a fast determination of the appropriate surfactant system. Also, as mentioned, not all surfactant systems lead to three-phase microemulsion systems but rather form gels or viscous phases over the salinity gradient, which is highly undesirable [11,12].

The study presented in this paper shows that a fast determination of the optimum formulation in petroleum system can be achieved by dynamic Salinity-Phase-Inversion (SPI) of emulsions. By analysing the SPI curves obtained, it is possible to identify the optimum salinity and to anticipate the formation of WIII systems or not at equilibrium. This technique significantly speeds up the screening of surfactant systems for EOR and other surfactant-based improved oil recovery (IOR) processes.

## 2. Material and methods

### 2.1. Chemicals

The two crude oils studied - noted A and B - were supplied by TOTAL SA. Their main characteristics such as TAN, TBN, ASCI index, API degree, SARA fractions, viscosity and density at 25°C are listed in Table 1. TAN and TBN were measured according to the standard methods ASTM D664 [13] and ASTM D2896 [14] respectively. The ASCI index was determined with a specific method developed by TOTAL [15]. API degree was calculated from density extrapolated at 15°C and SARA fractions were determined by TLC-FID [16]. The EOR surfactants used in this study were developed and supplied by BASF. Two families of surfactants have been investigated: sulfated alkoxy *n*-alcohols (*n*-C<sub>16-18</sub>PO<sub>k</sub>EO<sub>j</sub>SO<sub>4</sub>Na) as main surfactants and ethoxylated *i*-alcohols (*i*-C<sub>i</sub>EO<sub>j</sub>) as co-surfactant. Their hydrophobic tails are either linear or branched and have between 10 and 18 carbons. The well-defined *n*-C<sub>10</sub>EO<sub>4</sub> surfactant was synthesized in the lab and distilled to obtain an ultra-pure sample (> 99%) [17]. Bis(2-ethylhexyl) sulfosuccinate sodium salt, AOT (96%), *n*-Octane (≥ 99%), *n*-Decane (≥ 99%) and Toluene (≥ 99%) were purchased from Sigma Aldrich.

**Table 1:** Characteristics of the crude oils studied

Crude oil	A	B
TAN (mg KOH/g) <sup>a</sup>	0.5	2.0
TBN (mg eq KOH/g) <sup>b</sup>	1.7	2.5
ASCI index <sup>c</sup>	17	17
Viscosity at 25°C (mPa.s)	15.3	82.5
Density (25°C)	0.87	0.91
API degree <sup>d</sup>	31	23
Saturates (wt%) <sup>e</sup>	59.7	42.4
Aromatics (wt%) <sup>e</sup>	28.5	42.5
Resins (wt%) <sup>e</sup>	9.1	11.4
Asphaltenes(wt%) <sup>e</sup>	2.7	3.7

<sup>a</sup> TAN = Total Acid Number

<sup>b</sup> TBN = Total Base Number

<sup>c</sup> ASCI =Asphaltene Solubility Class Index

<sup>d</sup> API = American Petroleum Institute

<sup>e</sup> SARA = Saturates, Aromatics, Resins, Asphaltenes

## **2.2. Experimental procedure**

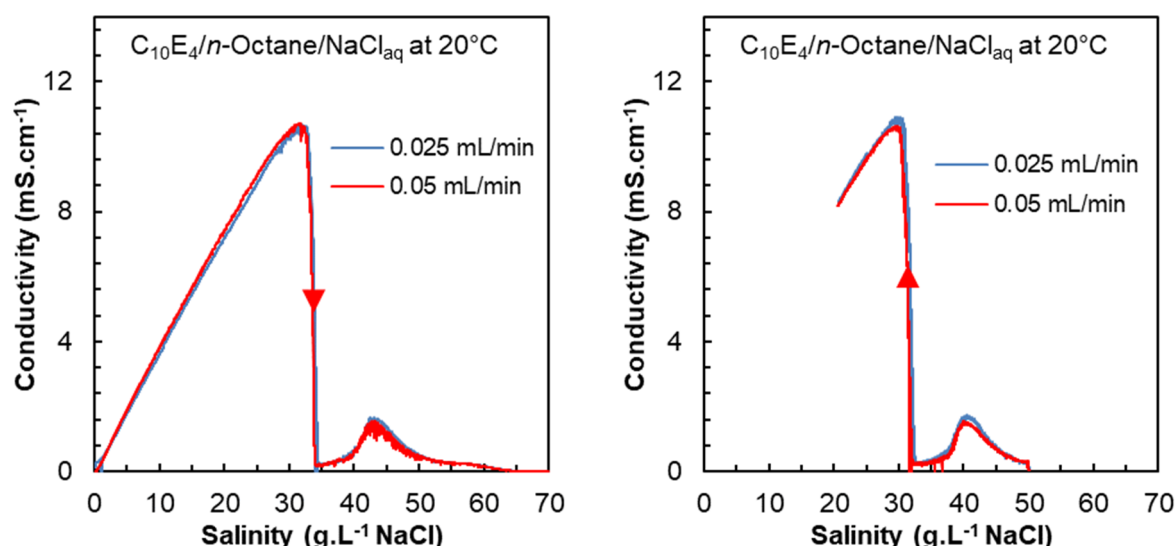
### **2.2.1. SOW equilibrium behaviour**

The phase behaviour of surfactants/oil/water systems was observed at WOR 1. Crude oils and model oils were used, and the water phase salinity was changed with NaCl with steps ranging from 1 g/L to 5 g/L depending on the systems. For crude oil-based systems, 1 mL of salted water containing the surfactant and the co-surfactant at the given concentration, with or without alkali (sodium carbonate or ammonia) was first poured in 5 mL-pipettes. Then, 1 mL of oil was added and the pipettes were sealed under a nitrogen flow and placed at the desired temperature for 10 minutes. They were then gently mixed and then mixed again after 1 hour, 2 hours and one night of equilibration. Observations of phase behaviour were made after complete stabilization (several days to several weeks). The optimal salinity was determined by plotting in the same graph the solubilisation of water in the microemulsion phase and the solubilisation of oil in the microemulsion phase, each divided by the total amount of surfactant and co-surfactant in the system, as a function of salinity [18,19]. The optimal salinity  $S^*$  is the salinity at which the curves cross. The maximum uncertainty is estimated at  $\pm 1$  g/L. The solubilisation ratio at optimum  $\sigma^*$ , defined in Eq. 5, is also determined at this point.

### **2.2.2. Dynamic Salinity Phase Inversion (SPI)**

Dynamic phase inversions were induced by increasing and decreasing continuously the aqueous phase salinity and were monitored by electrical conductivity measurement. Conductivity was recorded using a CDM210 conductivity meter from MeterLab® with a coupled conductivity-temperature electrode CDC641T from Radiometer Analytical®. Conductivity data were processed with the Labview software which records at the same time conductivity, temperature and the experimental time. The volume of the Surfactant/Oil/Water (SOW) mixtures varied during the experiment starting at 10 mL and finishing at 20 mL and the Water-to-Oil Ratio (WOR) was maintained equal to 1 (except for SPI experiments with  $C_{10}EO_4$  and AOT in which the water mass fraction  $f_w$  is maintained to 0.5). The emulsification vessel was a 2.5 cm diameter \* 20 cm height double-jacketed cylindrical tube. A 2 cm-

cross magnetic stirrer was used for agitation. The magnetic stirring rate was fixed at 900 rpm, which allowed obtaining reliable conductivity measurements in our systems, and the temperature was kept constant throughout the experiment by circulating water controlled by a HUBER 125 Ministat. To continuously modify the salinity, an aqueous solution containing both the surfactant(s) and concentrated or diluted NaCl was added to the initial SOW mixture at a controlled flow rate  $Q_i$  of 0.05 mL/min thanks to a press-syringe engines model 78-8100INT from KdScientific® fitted with a 10-mL Terumo-syringe (reference SS+10ES1). Simultaneously, the crude oil was introduced in the same way at the same flow rate in order to maintain a constant WOR. Using a flow rate of 0.05 mL/min is a good compromise between the length of the experiment and a low impact of salinity variation kinetics on the determination of emulsion conductivity, as shown in Figure 1.



**Figure 1:** Impact of flow rate on the conductivity profile during a dynamic SPI experiment with an increase (left) and a decrease (right) of the aqueous phase salinity for the model system 3 wt.%  $C_{10}EO_4/n$ -Octane/ $NaCl_{aq}$  system at 20°C at  $f_w = 0.5$ . The data have been recorded with a stirring rate of 900 rpm

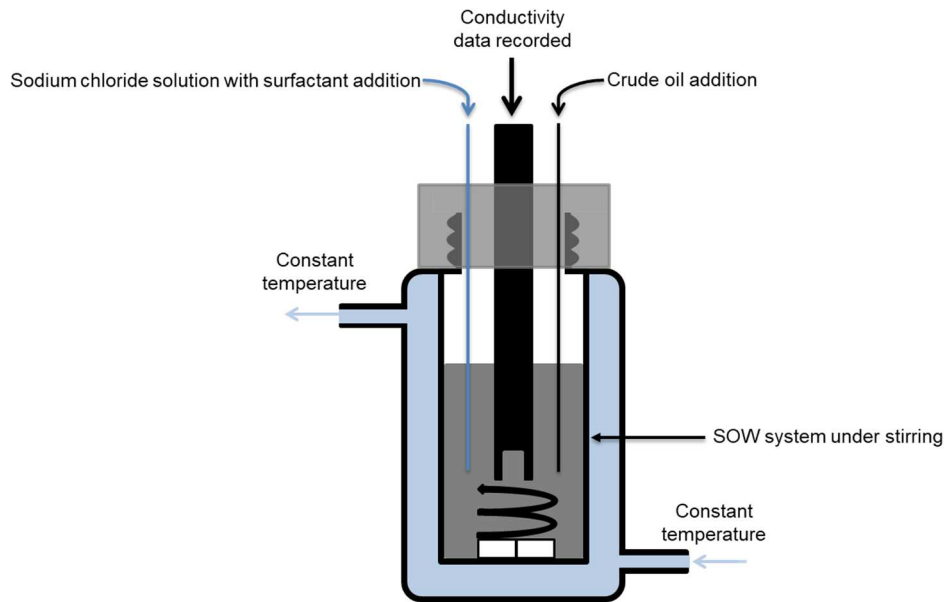
The procedure allows increasing or decreasing the aqueous phase salinity of the SOW sample under stirring without modifying the WOR nor the surfactant concentration, as illustrated in figure 2. Increasing the salinity of the aqueous phase requires dissolving large amounts of salt in an aqueous solution of surfactants, some of them being poorly salt-tolerant. Alternatively, two aqueous solutions

have to be injected, one containing the surfactant in low salinity brine, and the other one containing only a high concentration of salt. For simplification, the reverse procedure was implemented in this work, *i.e.* salinity was continuously decreased by adding a salt-free solution of surfactant to an initial SOW system with a high NaCl concentration. Such an experiment can be achieved within 30 minutes.

The aqueous phase salinity  $S(t)$ , expressed in grams of NaCl per liter of aqueous phase, can be calculated at each time of the experiment by the following relation:

$$S(t) = \frac{S_0 V_0 + t \cdot S_i Q_i}{V_0 + t \cdot Q_i} \quad (1)$$

where  $S_0$  and  $S_i$  are the initial and the added aqueous phase salinity respectively,  $t$  (in minutes) is the time,  $V_0$  (in mL) the initial aqueous volume and  $Q_i$  (in mL.min<sup>-1</sup>) the flow-rate of aqueous solution added to the SOW system. The SPI values are determined from the conductivity curves by using the parallel tangent methods and are given with a precision of  $\pm 0.5$  g.L<sup>-1</sup>.



**Figure 2:** Experimental set up to achieve the dynamic SPI of Surfactant/Crude oil/Brine systems by increasing or decreasing the aqueous phase salinity



### 2.2.3. Interfacial tension

The interfacial tensions were measured with a spinning drop tensiometer SDT from KRÜSS GmbH® allowing measurements down to  $10^{-6}$  mN.m<sup>-1</sup>. The measurements were carried out on equilibrated SOW systems at 55 °C and at different fixed aqueous phase salinities. The various phases (aqueous, oil and microemulsion phases) were first separated and were kept at 55°C during one hour before interfacial tension measurements. For SOW systems exhibiting WI and WII microemulsion behaviours at equilibrium, a droplet of the oil phase (between 2 and 10 µL) was inserted in the capillary filled with 1 mL of the aqueous phase. For WIII microemulsion systems, a 2 µL droplet of the oil phase in excess was inserted with a small amount of the middle-phase microemulsion (less than 1 µL in average) within the capillary filled with 1 mL of the aqueous phase in excess in order to get stable and accurate interfacial tension values as recommended in the literature [20]. All interfacial tension measurements were made after complete stabilisation of the SOW system inside the capillary and repeated on more than 5 droplets by using the Young-Laplace equation (Eq. 2).

$$\Delta P = 2\gamma H = \gamma \left( \frac{1}{R_1} + \frac{1}{R_2} \right) \quad (2)$$

where  $\Delta P$  is the Laplace pressure difference across the interface,  $\gamma$  the interfacial tension,  $H$  the mean curvature, and  $R_1$  and  $R_2$  are the principal radii of curvature [21]. For each individual droplet, the interfacial tension values are measured with a precision of  $\pm 1.10^{-6}$  mN.m<sup>-1</sup> according to the supplier. The values given are means values on 5 droplets and the mean deviation is 30% for WIII systems and 20% for WI and WII systems in average.

## 3. Results and discussion

In this section, after a description of the different detection methods, we show how the SPI method allows characterizing model SOW systems. Then, it is applied to petroleum SOW systems and we show the close correspondence between the SPI and the optimal salinity determined at equilibrium. We also demonstrate that the SPI method allows anticipating the formation of viscous phases.

194

### 195       **3.1. Detection methods of the phase inversion**

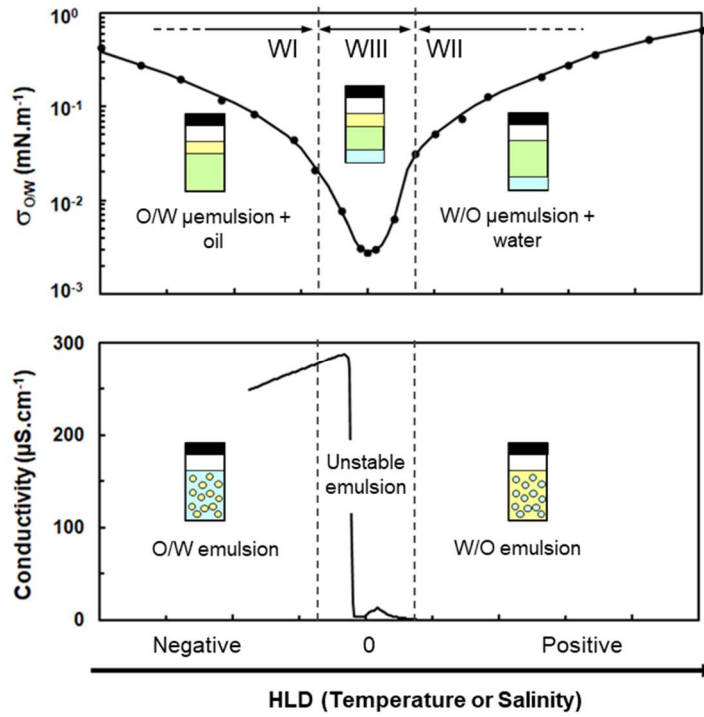
196   The so-called “standard” phase inversion is traditionally detected by emulsifying a series of tubes  
197   containing a SOW system pre-equilibrated at various temperatures or salinities [22]. The type of  
198   emulsion is then determined by conductimetry to identify the inversion border. It can also be  
199   determined in a dynamic way by applying a continuous variation of one of the formulation parameters  
200   to a stirred system. Dynamic experiments are usually performed using temperature as the formulation  
201   variable because it can easily and reversibly be changed in a controlled manner. The easiest way to  
202   determine the dynamic phase inversion transition of emulsions is to record the conductivity signals  
203   [23–33]. At the phase inversion, the SOW system reverses itself by giving a complex mixture of  
204   microemulsion, oil and water, which is less conductive than the previous O/W emulsion. It induces a  
205   fall in conductivity. After the phase inversion, the SOW system shows a conductivity close to zero  
206   because a W/O emulsion is obtained. The phase inversion is then defined as the temperature at which  
207   the fall in conductivity occurs. The dynamic method provides significant time savings compared to the  
208   equilibrium scan, a dramatic signal visualization of the phase inversion and a margin of error close to  
209   zero compared to the discontinuous equilibrium procedure with a series of pipettes [27]. The phase  
210   inversion can also be detected by viscosity measurements [34–38] or by light backscattering and near-  
211   infrared spectroscopy [39,40], as the light backscattering intensity strongly depends on the droplet size  
212   of the stirred emulsion [41,42].

213

### 214       **3.2. The Dynamic Salinity Phase Inversion (SPI) with model oils and surfactants**

215   The transitional phase inversion is defined as the change in the emulsion morphology from an oil-in-  
216   water (O/W) to a water-in-oil (W/O) emulsion, and vice versa, resulting from the change of the  
217   surfactant affinity from water to oil. This affinity change can be induced by modifying any  
218   formulation variable. Historically, temperature was used for polyethoxylated surfactant-based systems  
219   [23,24,43] but other variables like the number or ethoxy groups or the length of the hydrophobic tail of  
220   the surfactant, the nature and the concentration of alcohol in the blend, as well as the salinity of the

aqueous phase can also be used as variable parameters [25,26,44]. In the same way, numerous studies have evidenced that the phase inversion, induced by temperature or salinity, occurs within the WIII zone at equilibrium for model oils [25,26,45–49] and also for crude oils by taking temperature as formulation variable [34,50]. This result is valid whatever the type of surfactant if the volume Water-to-Oil Ratio (WOR) of the SOW system is close to 1 as shown in the literature for ionic [26] and nonionic [27,51,52] surfactants. This result is also valid even for mixtures of technical grade ionic and nonionic surfactants with variations of the proportion of surfactants in the mixture, as demonstrated by Antón *et al.* [53]. In addition to the WOR, other parameters can impact the identification of the optimum formulation by dynamic phase inversion. In particular, a too-low surfactant concentration, an inefficient stirring as well as a very high viscosity of the oil can impact the phase inversion conductivity signal and induce hysteresis (different inversion location depending on the way the formulation variable is changed), as highlighted by Márquez *et al.* in Temperature-induced phase inversion of non-ionic surfactants-based systems [54]. It means that, for suitable WOR, surfactant concentration, stirring and oil viscosity ranges, the phase inversion can be used to identify rapidly the optimum formulation for model oils (figure 3) [55] and also for crude oils when temperature is changed in a dynamic way [34,50].



**Figure 3:** Characteristic phenomena occurring close to the “optimum formulation” for (top) Equilibrated system: deep minimum of W/O interfacial tension and three-phase system with a middle microemulsion phase containing the same amount of oil and water and (down) Emulsified system: O/W, unstable, or W/O emulsion depending on whether the system is emulsified before, at, or after the optimum formulation and phase inversion of the same stirred system detected by a sudden drop of conductivity. Reproduced with authorization from Ref [55]

In a first step, the phase inversion induced by continuous salinity change was studied for two SOW model systems: the nonionic  $C_{10}EO_4/n$ -Octane/ $NaCl_{(aq)}$  and the ionic AOT/ $n$ -Decane/ $NaCl_{(aq)}$ .

For these simple SOW systems, the optimum salinities  $S^*$  can be approximately estimated with the Hydrophilic Lipophilic Deviation (HLD) equations from Salager *et al* (Eq. 3 for nonionics and 4 for ionics) [22].

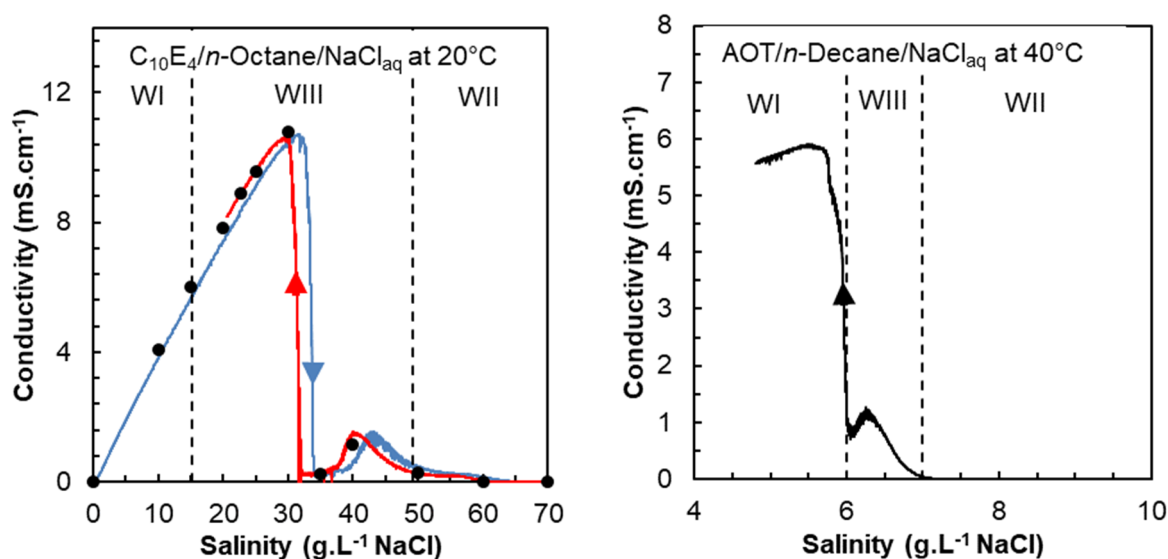
$$HLD = (\alpha - EON) - k \cdot EACN + t \cdot \Delta T + b \cdot S^* + f(A) = 0 \text{ at optimum formulation} \quad (3)$$

$$HLD = \sigma - k \cdot EACN + t \cdot \Delta T + \ln(S^*) + f(A) = 0 \text{ at optimum formulation} \quad (4)$$

Where:

- $\alpha$ ,  $\sigma$ , EON,  $k$ , and  $t$  are parameters depending on the surfactant,
- $S$  (wt%) is the salinity of the aqueous phase,
- EACN is the equivalent alkane carbon number of the oil,
- $\Delta T$  ( $^{\circ}\text{C}$ ) is the temperature deviation from  $25^{\circ}\text{C}$ ,
- $A$  (wt%) is the alcohol concentration and  $f(A)$  a function depending on the alcohol type.

The HLD relation establishes that the optimal salinity  $S^*$  and the optimal temperature are correlated in a logarithmic scale for ionic surfactants and in a linear way for nonionics. These equations predict that an optimal salinity of  $\approx 22 \text{ g.L}^{-1}$  at  $20^{\circ}\text{C}$  for the nonionic system and  $\approx 6 \text{ g.L}^{-1}$  at  $40^{\circ}\text{C}$  for the ionic system are required to reach the optimum formulation (calculations are detailed in the supplementary information) [22,33,56]. These systems were studied by continuously modifying the salinity to trigger the phase inversion. Figure 4 shows the conductivity vs salinity profiles for the nonionic (left) and ionic (right) systems. The vertical dotted lines in the figure indicate the frontiers of the different phase behaviors at equilibrium (WI, WIII and WII). For the 3 wt. %  $\text{C}_{10}\text{EO}_4/n\text{-Octane/NaCl}_{(\text{aq})}$  system, the inversion was studied both in dynamic (solid curves) and standard (dotted curve) mode. For the dynamic inversion, conductivity was monitored either by continuously increasing or decreasing the aqueous salinity (blue and red curves respectively in figure 4 left). For the standard inversion mode, the conductivity was measured after emulsification of pre-equilibrated samples at different salinities (black points in figure 4 left).



**Figure 4:** (Left) SPI of the 3 wt.%  $C_{10}EO_4/n$ -Octane/ $NaCl_{aq}$  system at  $20^\circ C$  with  $f_w = 0.5$  monitored by conductivity by continuously increasing (blue curve) or decreasing (red curve) the salinity, and after mixing samples initially at equilibrium (black dots) (Right) SPI of the 5 wt.% AOT/ $n$ -Decane/ $NaCl_{aq}$  system at  $40^\circ C$  with  $f_w = 0.5$  monitored in conductivity by decreasing the aqueous salinity. The equilibrium phase behavior (WI, WII or WIII) zones at different salinities are indicated in both figures (black dotted vertical lines).

As displayed in figure 4 (left), the SPI occurs in the salinity range  $31\text{--}34\text{ g.L}^{-1}\text{ NaCl}$  depending on the protocol used. This range value is somewhat higher than the value anticipated by the HLD equation. However, the dynamic SPI occurs within the WIII region, showing the effectiveness of this technique which can be used to identify rapidly the optimum formulation conditions of SOW systems based on nonionic surfactants. The profile of the blue curve follows the expected tendency, *i.e.* an increase of conductivity almost proportional to the NaCl concentration until a sharp drop when the morphology of the emulsion changes from O/W to W/O. The conductivity signal shows non-zero values between  $35$  and  $60\text{ g.L}^{-1}\text{ NaCl}$  with a slight maximum centered at  $43\text{ g.L}^{-1}\text{ NaCl}$ . This bump appears regardless the protocol used (continuous or discontinuous scan), indicating that it is not due to artefacts but results from the complex morphology of the three-phase emulsion Water/Microemulsion/Oil formed by agitation of the WIII system.

The  $S^*$  values determined by increasing or decreasing the salinity differ from 3 g.L<sup>-1</sup> NaCl. This small hysteresis could be due to the deviation from WOR = 1 when brine or pure water is added to the emulsion since the WOR is known to slightly influence the transitional phase inversion of emulsions [27]. For the same reason, the optimal salinity found by the discontinuous scan (black dots) differs slightly from that obtained under dynamic conditions. The dynamic SPI can be also used to identify the optimum salinity for ionic surfactants as demonstrated by figure 4 (right) for the 5 wt.% AOT/*n*-Decane/NaCl<sub>(aq)</sub> system at 40°C. For this system, the conductivity drop and the residual conductivity detected after the inversion coincides exactly with the WIII region observed at equilibrium, which confirms the close correspondence between the optimum formulation at equilibrium and the continuous SPI. The experimental SPI value of 6 g.L<sup>-1</sup> NaCl matches very well with the value predicted from the HLD equation but it slightly differs from the optimal salinity (7 to 9 g.L<sup>-1</sup>) reported by Kahlweit *et al.* for the same system [57]. Indeed, several artefacts can modify the experimental value of  $S^*$  especially when the WIII domain is narrow as for the AOT/*n*-Decane/NaCl<sub>(aq)</sub> system. Firstly, the gradual addition of pure water (decreasing salinity) to the initial SOW system changes a little bit the surfactant concentration (see figure 1 in supplementary information). Secondly, AOT (purity 96%) is a diester which may contain small amounts of the starting material (ethylhexyl alcohol) and can undergo some hydrolysis during storage. It is well known that even a small amount of this branched alcohol can play the role of co-surfactant and significantly shift the value of  $S^*$  [22,58,59].

### 3.3. The dynamic SPI for EOR applications

#### 3.3.1. Correspondence between phase behavior, emulsion inversion and interfacial tension minimum

The addition of well-chosen surfactants modifies the phase behavior of crude oil/water system and may eventually lead to the formation of three-phase microemulsion systems (WIII) in which the oil/water interfacial tension reaches an ultra-low value [60,20,22]. Such systems can be obtained by

balancing the surfactant affinity for the aqueous and oil phases to suit the crude oil under study. As already mentioned, the modification of the surfactant affinity can be made by adding a suitable co-surfactant or by tuning the temperature and/or the salinity [61].

In the EOR field, temperature is fixed by the reservoir temperature, and so salinity remains the preferred variable parameter to adjust the surfactant affinity. In that way, the optimum formulation is determined by discontinuous salinity scans carried out at equilibrium and at the reservoir temperature. From this salinity scan at equilibrium, it is possible to determine the optimum salinity  $S^*$  but also the efficiency of the surfactant system, which is evidenced by the microemulsion volume related to the volume of surfactant used. To express this efficiency, the solubilization ratio  $\sigma^*$ , expressed in  $\text{cm}^3/\text{cm}^3$  or cc/cc, is often used (Eq. 5) [4,62].

$$\sigma^* = \frac{V_{O \text{ or } W/\mu}}{V_S} = \frac{\text{Volume of water or oil in the microemulsion at optimum}}{\text{Volume of surfactant + co-surfactant in the system}} \quad (5)$$

From the solubilization ratio, it is also possible to infer the minimum interfacial tension at the optimum formulation by considering the model proposed by Huh displayed in Eq. 6 [18]:

$$\frac{\sigma^{*2} \cdot \gamma^*}{\cos(\frac{\pi}{4})} = a_H \quad (6)$$

where  $\sigma^*$  (cc/cc) is the solubilization ratio,  $\gamma^*$  ( $\text{mN} \cdot \text{m}^{-1}$ ) is the minimum of interfacial tension, and  $a_H$  ( $\text{mN} \cdot \text{m}^{-1}$ ) is a parameter that depends on the surfactant type.

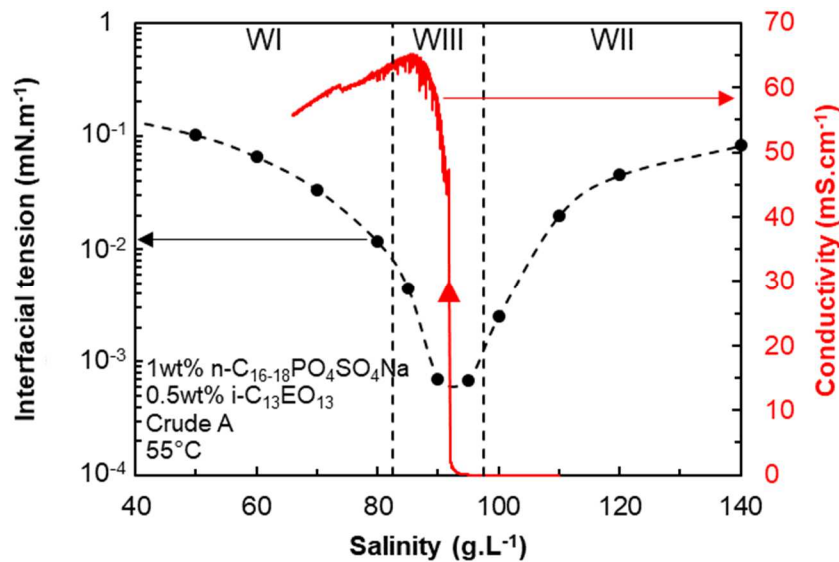
$a_H$  has already been calculated for series of surfactants [63–65] and can be approximated to 0.42  $\text{mN} \cdot \text{m}^{-1}$  in most cases. Therefore, the Huh's equation is often used in its simplified form (Eq. 7) [66]:



$$\gamma^* = \frac{0.3}{\sigma^{*2}} \quad (7)$$

Solubilization ratios higher than 10 cc/cc are usually desired for EOR applications, to ensure that sufficiently low interfacial tension values are attained.

As the dynamic SPI method allows finding the “optimal conditions” for model systems, this technique deserves to be tested on systems relevant to EOR applications, for which the optimum salinity is a key parameter. In order to evaluate the feasibility of the SPI with petroleum-based systems, a mixture of a sulfated alkoxyated alcohol surfactant, a nonionic co-surfactant, a typical crude oil and brine was studied. Figure 5 shows the conductivity vs. salinity profile for the  $[n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na} + i\text{-C}_{13}\text{EO}_{13}]$ /Crude oil A/Brine at WOR=1. Additionally, the O/W interfacial tensions measured on equilibrated systems with a spinning-drop tensiometer at different salinities are also shown.



**Figure 5:** Identification of the “optimum salinity”  $S^*$  of a crude oil based SOW system by dynamic SPI monitored by conductimetry while decreasing the aqueous salinity (red curve). Evolution of the W/O interfacial tension (black dots and dashed curve) and limits of the WIII region (black dotted vertical lines)

363

364 As can be seen in figure 5, the dynamic SPI coincides exactly with the minimum interfacial tension.

365 As for model SOW system, the sharp drop in conductivity occurs within the salinity range where the

366 WIII microemulsion system at equilibrium is obtained with ultra-low interfacial tension values close

367 to  $5.10^{-4}$  mN.m<sup>-1</sup>. By comparison, the Huh equation gives a value of  $2.5.10^{-3}$  mN.m<sup>-1</sup> for this system

368 according to the solubilization ratio at equilibrium (11 cc/cc). This predicted value is significantly

369 higher than the measured IFT value, which indicates that the  $a_H$  parameter commonly taken in the

370 Huh's model overestimates the interfacial tension value at optimum for this surfactant system.

371 Anyway, this experiment highlights that the SPI actually occurs at the optimum formulation, even for

372 SOW systems composed of crude oils and mixtures of ionic/nonionic technical grade surfactants.

373 The evolution of the interfacial tension appears to be similar to the one reported in literature using

374 salinity as formulation variable for ionic surfactants [67,68] and temperature [20] or the number of

375 ethylene oxide groups for nonionic surfactants [69].

376

### 377 **3.3.2. Effect of formulation on the formation of viscous phases for equilibrated SOW**

#### 378 **systems**

379 One difficulty often encountered when formulating surfactant blends for EOR is that some crude oils

380 form undesirable viscous phases instead of the fluid WIII systems. To overcome this problem, the

381 formulation must be modified. In practice, an alkali is often used to reveal the endogenous surfactants

382 and/or significant amounts of a co-solvent are added to the formulation in order to make the interfacial

383 film more fluid [70–72]. Barnes *et al.* have shown that a given surfactant formulation matches well,

384 moderately or badly the oil (*i.e.* it forms WIII systems, loose or viscous gel phases respectively)

385 depending on the crude chemical nature and they have tried to relate these behaviours with some bulk

386 analytical descriptors of the oil [12]. Crude oils also contain naphthenic acids and asphaltenes [11]

387 acting as endogenous surfactants that play a role in the SOW phase behaviour in the presence of

synthetic surfactants [73]. This effect is particularly sensitive for high TAN crude oils in the presence of alkali because naphthenic acids are converted into highly surface active naphthenates.

Table 2 shows some surfactant formulations studied at equilibrium with crude A (entries a-j) and crude B (entries k-r). They are all based on a mixture of an ionic surfactant  $n\text{-C}_{16-18}\text{PO}_k\text{EO}_j\text{SO}_4\text{Na}$  and a nonionic co-surfactant ( $i\text{-C}_i\text{EO}_j$ ) or 2-Butanol, with or without alkali (sodium carbonate or ammonia). This set of experiments has been built starting from the conditions described in figure 5 with crude A (entry a). The initial formulation has been changed to study the influence of the formulation variables on the optimal salinities  $S^*$  and on the efficiencies of the surfactant systems (solubilisation ratio values  $\sigma^*$  or formation of viscous phases). These effects are summarized in Table 3.

**Table 2:** Experimental values obtained for optimum salinity at equilibrium  $S^*$  and solubilization ratio at optimum  $\sigma^*$  for the different petroleum SOW systems tested

Entry	Crude	Ionic surfactant		Co-surfactant		Alkali		T °C	$S^*$ g.L <sup>-1</sup>	$\sigma^*$ cc/cc
		Formula	wt. %	Formula	wt. %	Type	wt. %			
a	A	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	-	-	55	89	11
b	A	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{10}\text{EO}_{10}$	0.33	-	-	55	60-70 <sup>§</sup>	N/A
c	A	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{10}\text{EO}_{10}$	0.5	-	-	55	85	10
d	A	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{10}\text{EO}_{10}$	1	-	-	55	110	5
e	A	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{10}$	0.5	-	-	55	55-70 <sup>§</sup>	N/A
f	A	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{17}\text{EO}_{12}$	0.5	-	-	55	50-70 <sup>§</sup>	N/A
g	A	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	2-BuOH	1.5	-	-	55	40-45 <sup>§</sup>	N/A
h	A	$n\text{-C}_{16-18}\text{PO}_2\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	-	-	65.5	81	9
		/ $n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$ (50/50)								
i	A	$n\text{-C}_{16-18}\text{PO}_2\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	$\text{Na}_2\text{CO}_3$	0.5	65.5	70	10
		/ $n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$ (50/50)								
j	A	$n\text{-C}_{16-18}\text{PO}_2\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	$\text{NH}_{3,\text{aq}}$	0.5	65.5	70	13
		/ $n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$ (50/50)								
k	B	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	$\text{Na}_2\text{CO}_3$	0.5	40	150	8
l	B	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	$\text{Na}_2\text{CO}_3$	0.5	55	134	8

m	B	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	$\text{Na}_2\text{CO}_3$	0.5	65	120	7
n	B	$n\text{-C}_{16-18}\text{PO}_7\text{EO}_{0.1}\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.25	$\text{Na}_2\text{CO}_3$	0.5	55	68	22
o	B	$n\text{-C}_{16-18}\text{PO}_7\text{EO}_{0.1}\text{SO}_4\text{Na}$	0.5	$i\text{-C}_{13}\text{EO}_{13}$	0.125	$\text{Na}_2\text{CO}_3$	0.5	55	79	20
p	B	$n\text{-C}_{16-18}\text{PO}_7\text{EO}_{0.1}\text{SO}_4\text{Na}$	0.25	$i\text{-C}_{13}\text{EO}_{13}$	0.0625	$\text{Na}_2\text{CO}_3$	0.5	55	81	24
q	B	$n\text{-C}_{16-18}\text{PO}_7\text{EO}_{0.1}\text{SO}_4\text{Na}$	0.25	$i\text{-C}_{13}\text{EO}_{13}$	0.0625	$\text{NH}_{3,\text{aq}}$	0.5	55	55	40
r	B +14% Toluene	$n\text{-C}_{16-18}\text{PO}_7\text{EO}_{0.1}\text{SO}_4\text{Na}$	0.25	$i\text{-C}_{13}\text{EO}_{13}$	0.0625	$\text{Na}_2\text{CO}_3$	0.5	55	55	30

<sup>§</sup>No VIII, gels formed instead in the salinity range indicated in the S\* column.

**Table 3: Effect of the main formulation variable changes on the phase behavior of the SOW systems described in Table 2**

Parameters	Entries	Formulation change	S*	$\sigma^*$
Amount of co-surfactant	b $\rightarrow$ c $\rightarrow$ d	% $i\text{-C}_{10}\text{EO}_{10}$ $\nearrow$	$\nearrow$	$\searrow$
Type of co-surfactant	a $\rightarrow$ c	$i\text{-C}_{13}\text{EO}_{13} \rightarrow i\text{-C}_{10}\text{EO}_{10}$	$\approx$	$\approx$
	c $\rightarrow$ e	$i\text{-C}_{10}\text{EO}_{10} \rightarrow i\text{-C}_{13}\text{EO}_{10}$	$\Rightarrow$ gel phase	NA
	c $\rightarrow$ f	$i\text{-C}_{10}\text{EO}_{10} \rightarrow i\text{-C}_{17}\text{EO}_{12}$	$\Rightarrow$ gel phase	NA
	c $\rightarrow$ g	$i\text{-C}_{10}\text{EO}_{10} \rightarrow 2\text{-BuOH}$	$\Rightarrow$ gel phase	NA
Base to reveal endogenous surfactants	h $\rightarrow$ i	+ $\text{Na}_2\text{CO}_3$ (crude A)	$\searrow$	$\approx$
	h $\rightarrow$ j	+ $\text{NH}_{3,\text{aq}}$ (crude A)	$\searrow$	$\nearrow$
	p $\rightarrow$ q	$\text{Na}_2\text{CO}_3 \rightarrow \text{NH}_{3,\text{aq}}$ (crude B)	$\searrow$	$\nearrow$
Temperature	k $\rightarrow$ l $\rightarrow$ m	Temperature $\nearrow$	$\searrow$	$\approx$
Amount of {surfactant + co-surfactant}	n $\rightarrow$ o $\rightarrow$ p	%{surfactant + co-surfactant} $\searrow$	$\nearrow$	$\approx$
Type of oil	p $\rightarrow$ r	% Toluene $\nearrow$	$\searrow$	$\nearrow$

#### Amount and type of co-surfactant:

For crude A, poor in naphthenic acids and asphaltenes, the addition of alkali is not essential to obtain a VIII system but the nature and the amount of nonionic co-surfactant has to be adapted to avoid the formation of gel phases. A ratio surfactant/co-surfactant of 2 is convenient when the co-surfactant is  $i\text{-C}_{13}\text{EO}_{13}$  (entry a) or  $i\text{-C}_{10}\text{EO}_{10}$  (entry c). In contrast, the less hydrophilic co-surfactants  $i\text{-C}_{13}\text{EO}_{10}$  (entry e) or  $i\text{-C}_{17}\text{EO}_{12}$  (entry f) [34] lead to the formation of gel phases. Increasing the amount of

nonionic co-surfactant  $i\text{-C}_{10}\text{EO}_{10}$  is a lever to overcome the formation of gel phases (see entries b-d) but it has a negative impact on the solubilisation ratio. Using a simple alcohol (2-butanol), even in large amounts, is not sufficient to avoid the formation of gel phases without alkali (entry g).

#### *Revelation of endogenous surfactants with alkali:*

Crude B contains a significant amount of naphthenic acids according to its TAN value (see Table 1). It is a more “difficult” oil than crude A because it is significantly more viscous and because it is particularly prone to the formation of gel phases. For crude oil B, the addition of an alkali is necessary to obtain WIII systems with these surfactants blends. The naphthenates formed at high pH probably act as anionic co-surfactants and promote the formation of fluid interfaces. On the contrary, crude A that has a relatively low TAN (see Table 1) does not require the systematic use on alkali to obtain WIII systems. The addition of either sodium carbonate or ammonia induces a slight decrease of the optimal salinity (entries h-j). For crude B, the impact of alkali is much more pronounced. First, a decrease of the total concentration of exogenous surfactant and cosurfactant at a constant  $\text{Na}_2\text{CO}_3$  concentration (entries n-p) induces a significant increase of the optimal salinity.

The increase in hydrophilic character of the surfactant blend could have several origins. Exogenous surfactants are not pure but technical grade compounds, and a change in concentration can lead to a change in the nature of the surfactants present at the interface (more hydrophilic in this case) due to different partition coefficients. Another hypothesis is the contribution of hydrophilic naphthenates because their relative amount (compared to exogenous surfactants) increases. It can also result from a combination of both effects. The second noticeable impact is the nature of the alkali. The addition of  $\text{NH}_{3,\text{aq}}$  make the surfactant blend much more hydrophobic (compare  $S^*$  for entries p and q), which can be due to different natures and/or amounts of the endogenous surfactants revealed. The values of solubilisation ratios obtained for crude B in the presence of alkali are only apparent and are overestimated since the naphthenates formed are not taken into account in the calculation.

#### *Extrapolation to Reservoir conditions:*

The SOW phase behaviour needs to be extrapolated to reservoir conditions. Temperature has an important effect on the optimal salinity (entries k-m) for these systems. An increase in temperature leads to a decrease of the optimum salinity because surfactants loose hydrophilicity, in agreement with previous studies with crude oils [34]. It does not impact the efficiency of the surfactant system, because of this negative salinity shift, as evidenced by the constant solubilisation ratios. To take into account the modification of the oil physical chemistry induced by the dissolution of gases in the (P,T) reservoir conditions, surrogate oils are usually prepared by blending the dead crude oil with a defined amount of solvent. Toluene is usually chosen because its EACN equals 1 and it is therefore a good substitute of methane at ambient pressure [74]. As expected, adding a significant amount of toluene to crude B in order to mimic the properties of the live oil induces a decrease of the optimal salinity and an increase of the solubilisation ratio (see entries p and r).

### 3.3.3. Identification of the formation of viscous phases by continuous SPI experiments

To evidence that the dynamic SPI can be an effective tool for optimizing surfactant blends for EOR, the SPI has been determined for a selection of the SOW systems described in Table 2 (entries a-g, k-m, o-p and r). Experimental results are gathered in Table 4.

**Table 4:** Petroleum-based SOW systems investigated by dynamic SPI. The SPI column shows the salinity values at which the phase inversion is observed. Columns  $S^*$  and  $\sigma^*$  correspond to the value determined for the same system at equilibrium

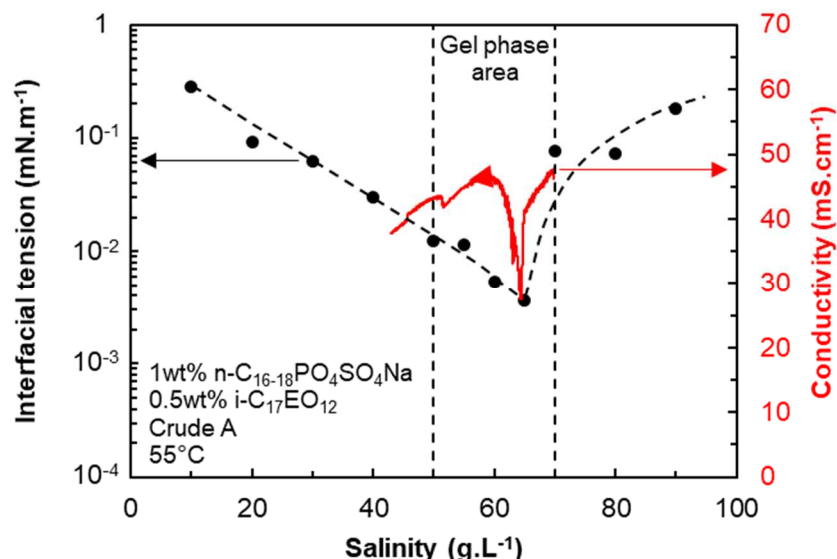
Entry	Crude	Ionic surfactant		Co-surfactant		Alkali		T °C	$S^*$ g.L <sup>-1</sup>	$\sigma^*$ cc/cc	SPI g.L <sup>-1</sup>
		Formula	wt.%	Formula	wt.%	Type	wt.%				
a	A	<i>n</i> -C <sub>16-18</sub> PO <sub>4</sub> SO <sub>4</sub> Na	1	<i>i</i> -C <sub>13</sub> EO <sub>13</sub>	0.5	-	-	55	89	11	91
b	A	<i>n</i> -C <sub>16-18</sub> PO <sub>4</sub> SO <sub>4</sub> Na	1	<i>i</i> -C <sub>10</sub> EO <sub>10</sub>	0.33	-	-	55	60-70 <sup>§</sup>	N/A	77
c	A	<i>n</i> -C <sub>16-18</sub> PO <sub>4</sub> SO <sub>4</sub> Na	1	<i>i</i> -C <sub>10</sub> EO <sub>10</sub>	0.5	-	-	55	85	10	91
d	A	<i>n</i> -C <sub>16-18</sub> PO <sub>4</sub> SO <sub>4</sub> Na	1	<i>i</i> -C <sub>10</sub> EO <sub>10</sub>	1	-	-	55	110	5	114
e	A	<i>n</i> -C <sub>16-18</sub> PO <sub>4</sub> SO <sub>4</sub> Na	1	<i>i</i> -C <sub>13</sub> EO <sub>10</sub>	0.5	-	-	55	55-70 <sup>§</sup>	N/A	73 <sup>§</sup>
f	A	<i>n</i> -C <sub>16-18</sub> PO <sub>4</sub> SO <sub>4</sub> Na	1	<i>i</i> -C <sub>17</sub> EO <sub>12</sub>	0.5	-	-	55	50-70 <sup>§</sup>	N/A	64 <sup>§</sup>
g	A	<i>n</i> -C <sub>16-18</sub> PO <sub>4</sub> SO <sub>4</sub> Na	1	2-BuOH	1.5	-	-	55	40-45 <sup>§</sup>	N/A	49 <sup>§</sup>

k	B	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	$\text{Na}_2\text{CO}_3$	0.5	40	150	8	150
l	B	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	$\text{Na}_2\text{CO}_3$	0.5	55	134	8	127
m	B	$n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$	1	$i\text{-C}_{13}\text{EO}_{13}$	0.5	$\text{Na}_2\text{CO}_3$	0.5	65	120	7	110
o	B	$n\text{-C}_{16-18}\text{PO}_7\text{EO}_{0.1}\text{SO}_4\text{Na}$	0.5	$i\text{-C}_{13}\text{EO}_{13}$	0.125	$\text{Na}_2\text{CO}_3$	0.5	55	79	20	80
p	B	$n\text{-C}_{16-18}\text{PO}_7\text{EO}_{0.1}\text{SO}_4\text{Na}$	0.25	$i\text{-C}_{13}\text{EO}_{13}$	0.0625	$\text{Na}_2\text{CO}_3$	0.5	55	81	24	85
r	B +14% Toluene	$n\text{-C}_{16-18}\text{PO}_7\text{EO}_{0.1}\text{SO}_4\text{Na}$	0.25	$i\text{-C}_{13}\text{EO}_{13}$	0.0625	$\text{Na}_2\text{CO}_3$	0.5	55	55	30	53

<sup>§</sup>No WIII, gels formed instead in the salinity range indicated in the S\* column. The SPI values are determined at the negative spike (see figure 6)

In a previous work, we have shown that there is a correspondence between the conductivity profile obtained during the dynamic PIT of petroleum systems and their ability to form WIII systems at equilibrium [34]. Figure 5 and Figure 6 show that it holds true when the inversion is induced by a continuous variation of salinity. The system based on  $n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$  1 % /  $i\text{-C}_{13}\text{EO}_{13}$  0.5 % (Table 4 entry a, figure 5) forms WIII systems at equilibrium and shows a complete inversion on the dynamic SPI conductivity profile. On the contrary, the system based on  $n\text{-C}_{16-18}\text{PO}_4\text{SO}_4\text{Na}$  1 % /  $i\text{-C}_{17}\text{EO}_{12}$  0.5 % (Table 4 entry f, figure 6) does not form WIII systems at equilibrium but rather gel phases on a certain range of salinities. During the dynamic SPI, no clean inversion is observed, however an accident on the conductivity curve is clearly visible. The same behavior occurs for all systems that do not form WIII systems at equilibrium (entries e-g of table 4) except for the system of entry b, which will be discussed in the following section.

The salinity at which the conductivity shows a negative spike is related to the zone of gel phase formation at equilibrium. For these systems, the oil/water interfacial tension is reduced when approaching the optimal salinity range ( $3.6 \cdot 10^{-2} \text{ mN.m}^{-1}$  measured for the system described in Table 4, entry f at 65 g/L), however, it does not reach ultra-low values. We can suppose that the co-surfactant chosen in this case is not suitable to promote a fluid interface allowing the inversion of the emulsion during the dynamic salinity scan and the formation of a WIII system at equilibrium.



**Figure 6:** Identification of the optimum salinity range of a SOW system exhibiting viscous phases at equilibrium (entry *f* of Table 4) by dynamic SPI monitored by conductimetry while decreasing the aqueous salinity (red curve). Evolution of the W/O interfacial tension (black dots and dashed curve) and gel phase area borders (back dotted vertical lines)

From a practical point of view, the shape of the conductivity profile during a dynamic SPI can be used as a descriptor of the phase behaviour of petroleum SOW systems regarding their ability to form a W/O microemulsion system or not.

### 3.3.4. Use of the dynamic SPI to quantify the influence of formulation variables on the position of the optimum

Also, the dynamic SPI reveals to be a valuable tool to quantify the influence of cross-parameters on the optimum formulation. Figures 7 and 8 illustrate the impact of temperature and co-surfactant concentration respectively.

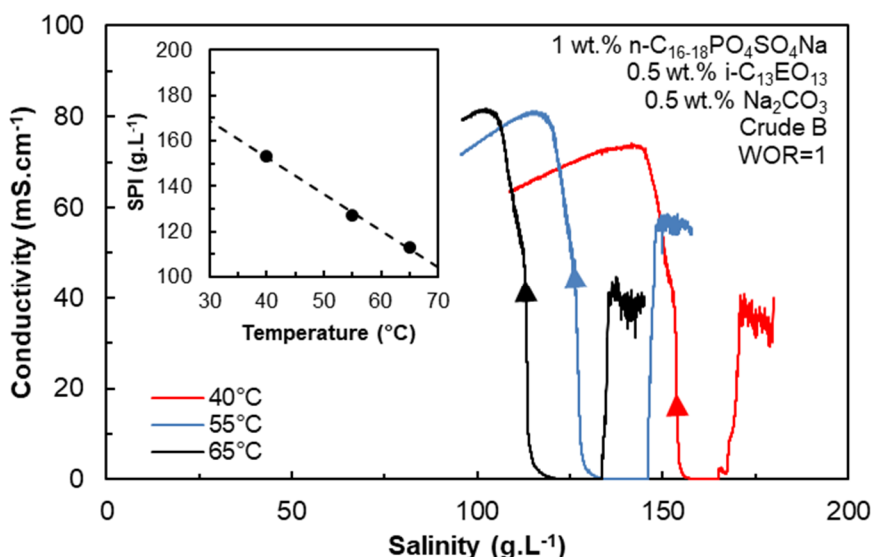
Figure 7 shows the conductivity profiles during the dynamic SPI of the *n*-C<sub>16-18</sub>PO<sub>4</sub>SO<sub>4</sub>Na 1 % / *i*-C<sub>13</sub>EO<sub>13</sub> 0.5 % with crude B in the presence of Na<sub>2</sub>CO<sub>3</sub> (Table 3, entries k-m). A temperature increase leads to a decrease of *S*\* at equilibrium and of the SPI under dynamic conditions. Increasing



temperature decreases the hydrophilicity of the polyethoxylated nonionic co-surfactant due to the progressive dehydration of the polyethoxylated polar head. The same effect occurs for the propoxylated groups of the hybrid anionic propoxylated sulfate, whereas the ionic head itself becomes more hydrophilic with increasing temperature.

The fact that the addition of propoxylated groups within a homogeneous series of ionic surfactants decrease the global hydrophilicity towards temperature means that propoxylated groups are more impacted by temperature changes than the ionic head [34,50]. As a consequence, the salinity needed to attain the optimum conditions is lower because the temperature increase has made the surfactant system more hydrophobic, as already mentioned by Hammond *et al.* with the same class of surfactants and model oils [75]. The linear evolution of the SPI value with temperature (insert in figure 7) indicates that the surfactant blend as a whole shows a nonionic-type dependence towards temperature (HLD Eq. 3).

It is important to stress that these dynamic SPI experiments have been performed by continuously decreasing the salinity, as indicated by the direction of the arrows on the conductivity profiles in Figure 7. Contrary to expectations, a noisy conductivity signal is always measured at high salinity, at the beginning of the experiment, whereas equilibrated systems show a regular WII behavior at these salinities. This phenomenon is due to a poor dispersion of droplets at the beginning of the emulsification, due to the mild stirring and high interfacial tensions, which does not allow to get the expected W/O emulsion. When the salinity approaches the optimal salinity range, the interfacial tension falls, which favors the dispersion of droplets into the expected W/O emulsion morphology, as indicated by the zero-conductivity signal. When salinity further decreases and reaches the optimal salinity, the conductivity sharply increases at the phase inversion towards an O/W morphology.

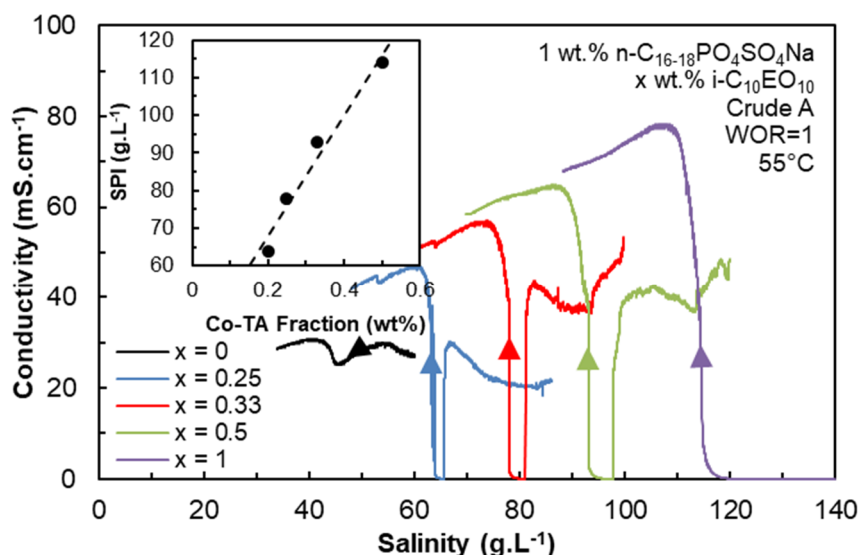


**Figure 7:** Influence of temperature on the dynamic SPI of a petroleum SOW system (entries k-m of Table 4)

As the experiments were performed only by decreasing continuously the salinity, we could not evidence hysteresis phenomena, but we can suspect they exist for this type of systems. As highlighted in Figure 2 in Supplementary information, this hysteresis is more pronounced for crude oil-based systems than for the model system 3 wt. % C<sub>10</sub>EO<sub>4</sub>/n-Octane/NaCl<sub>(aq)</sub> previously studied. However, it does not impede the identification of the SPI even if the viscosity of the crude oil is quite high (176 mPa.s at 25°C). Figure 2 in SI also demonstrates that keeping a constant volume during the experiment can increase the precision in the identification of the optimum formulation.

Figure 8 illustrates the impact of the concentration of nonionic co-surfactant for the *n*-C<sub>16-18</sub>PO<sub>4</sub>SO<sub>4</sub>Na 1 % / *i*-C<sub>10</sub>EO<sub>10</sub> x % with crude A without alkali (Table 4, entries b-d). Increasing the co-surfactant concentration increases the global hydrophilicity of the surfactant system, as indicated by the higher salinity required to attain the optimum formulation. A similar trend has already been reported by *Salager et al.* who showed that increasing the amount of ethoxylated nonionic surfactant in a nonionic/ionic mixture leads to an increase of optimum salinities [76]. For this system, the dynamic SPI is linearly correlated with the amount of nonionic co-surfactant in the blend, as indicated by the insert in Figure 8.

In this case also, it is interesting to look at the shapes of the conductivity profiles closely. Again, the salinity has been continuously decreased to pass through the conditions of phase inversion. For the system having the highest co-surfactant content (and therefore also the highest total surfactant concentration), the expected W/O emulsion is obtained at high salinity at the beginning of the experiment, as indicated by the zero-conductivity signal (violet curve in Figure 8). When the co-surfactant content is reduced (green, red and blue curves), the same phenomenon as the one described in Figure 8 is observed: the total amount of surfactant and/or the content of co-surfactant are not enough to stabilize properly water droplets in oil at high salinity at the beginning of the experiment (non-zero conductivity). When the salinity approaches the SPI, the oil/water interfacial tension is reduced enough to favor the dispersion and a zero-conductivity is measured. The emulsion then reverts to an O/W emulsion of high conductivity at the SPI. The range in which a W/O emulsion (zero-conductivity signal) is obtained is very reduced for the system with 0.33% co-surfactant (red curve in Figure 8), and even more for the system containing 0.25% co-surfactant (blue curve in Figure 8) which do not form WIII systems at equilibrium (Table 2, entry o). For these concentrations, the shape of the conductivity curves during the dynamic SPI gets close to what was observed in Figure 6, indicating a very narrow area of low interfacial tensions (less than 3 g.L<sup>-1</sup>). A non-inversion is observed when no co-surfactant is added to the formulation (black curve in Figure 8).

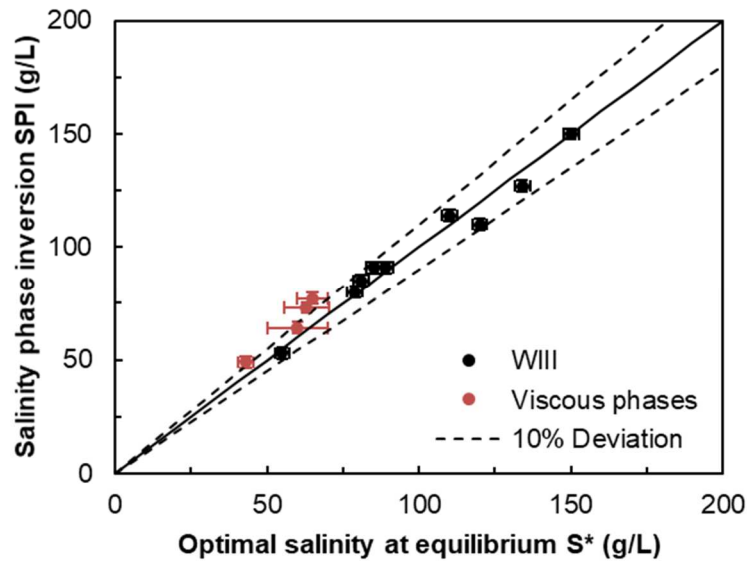


**Figure 8:** Influence of the nonionic co-surfactant concentration on the SPI of a petroleum SOW system (Table 4, entries b-d)

### 3.3.5. Accuracy of the determination of optimal salinity by dynamic SPI

Finally, the data presented in Table 4 are represented graphically in Figure 9 to highlight the good correspondence between the dynamic SPI and the optimal salinity  $S^*$  determined at equilibrium. A deviation lower than 10 % is observed. These data validate the dynamic SPI method as an alternative to the traditional static scans to find the conditions to reach the optimum formulation.

Its main advantage is a rapid execution - thirty minutes in average - coupled with a limited use of reactants. In the case of systems giving viscous phases instead of WIII microemulsions, a “SPI” is taken at the minimum of conductivity (see figure 6) and correlates with the salinity range in which gel phases are formed at equilibrium.



**Figure 9:** Correlation between the dynamic salinity of phase inversion SPI and the optimal salinity determined at equilibrium  $S^*$  for the different petroleum-based SOW systems presented in Table 4. In the case of non-inversion, the SPI is taken at the salinity of the negative spike (see Figure 6) and it is related to the salinity range of formation of gel phases at equilibrium

## Conclusions

In this work, a new method to achieve a fast detection of the optimum formulation of Surfactant/Crude oil/Brine at WOR in the vicinity of 1 is described. It is based on a continuous modification of the salinity of a stirred and thermostated SOW system to cause a dynamic Salinity Phase Inversion (SPI) of the emulsion detected by a sudden jump in conductivity. The study demonstrates that the phase inversions of model and petroleum SOW systems are closely linked to their phase behavior at equilibrium. The SPI experiment can be used to determine optimal salinities in a record time of thirty minutes compared to equilibrium salinity scans carried out in the conventional manner. Moreover, the shape of the conductivity curve was found to be a relevant descriptor for predicting and preventing the formation of viscous phases generated when inappropriate surfactant blends are used with a given crude oil.

For all these reasons, the dynamic SPI technique is particularly suitable for EOR applications and thanks to its fast implementation, it is particularly efficient for screening a large number of formulations in order to find rapidly the most effective surfactant systems. A variation of the technique is to trigger the phase inversion by continuously modifying the surfactant blend at a constant salinity. These dynamic experimental techniques can be carried out in conjunction with predictive models already described in the literature to accelerate even more the surfactant formulation screening [77–83].

More generally, this method has also a great potential for quantifying the sensitivity of surfactants to electrolytes. This will be described in a forthcoming paper. Finally, the SPI technique could be advantageously implemented to select the most efficient surface-active additives for other crude oil processing operations, such as wellbore cleaning, well injectivity restoration or demulsification.

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## Conflict of interest

The authors declare that they have no conflict of interest.

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