Interfacial and Colloidal Forces Governing Oil Droplet Displacement: Implications for Enhanced Oil Recovery

Suparit Tangparitkul 1,2, Thibaut V. J. Charpentier 1, Diego Pradilla 3 and David Harbottle 1,*

1 School of Chemical and Process Engineering, University of Leeds, Leeds LS2 9JT, UK; suparit.t@cmu.ac.th (S.T.); t.charpentier@leeds.ac.uk (T.V.J.C.)
2 Department of Mining and Petroleum Engineering, Chiang Mai University, Chiang Mai 50200, Thailand
3 Grupo de Diseño de Producto y de Proceso (GDPP), Departamento de Ingeniería Química, Universidad de los Andes, Carrera 1 este No. 18A-12, Edificio Mario Laserna, Piso 7, Bogotá 111711, Colombia; d-pradil@uniandes.edu.co
* Correspondence: d.harbottle@leeds.ac.uk; Tel.: +44-113-343-4154

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Abstract: Growing oil demand and the gradual depletion of conventional oil reserves by primary extraction has highlighted the need for enhanced oil recovery techniques to increase the potential of existing reservoirs and facilitate the recovery of more complex unconventional oils. This paper describes the interfacial and colloidal forces governing oil film displacement from solid surfaces. Direct contact of oil with the reservoir rock transforms the solid surface from a water-wet to neutrally-wet and oil-wet as a result of the deposition of polar components of the crude oil, with lower oil recovery from oil-wet reservoirs. To enhance oil recovery, chemicals can be added to the injection water to modify the oil-water interfacial tension and solid-oil-water three-phase contact angle. In the presence of certain surfactants and nanoparticles, a ruptured oil film will dewet to a new equilibrium contact angle, reducing the work of adhesion to detach an oil droplet from the solid surface. Dynamics of contact-line displacement are considered and the effect of surface active agents on enhancing oil displacement discussed. The paper is intended to provide an overview of the interfacial and colloidal forces controlling the process of oil film displacement and droplet detachment for enhanced oil recovery. A comprehensive summary of chemicals tested is provided.

Keywords: enhanced oil recovery; oil film displacement; colloid and interfacial science; wettability; surfactants; nanoparticle fluids

1. Introduction

The global energy landscape is gradually transitioning towards renewables, however, a reliance on non-renewables, particularly petroleum, will remain for several decades due to its importance as a fuel and chemical feedstock, which is a critical component to the steady improvement in the quality of life of developing countries. While developed countries take the lead on demonstrating the application of non-renewables, their remaining reliance on petroleum as part of the energy matrix remains for the foreseeable future (beyond 2050) [1]. With overall petroleum demand expected to increase [2], demand can only be met by increasing global production, which also coincides with the depletion of ‘easy-to-produce’ oil.

With few giant oil fields being discovered and new reserves frequently identified in remote/challenging locations, there is a growing need to increase the potential of existing reserves and improve the worldwide average oil recovery factor from as low as 20% to 40%. Methods of enhanced
oil recovery (EOR) have the potential to double the produced lifetime of existing proven reserves; which have a current lifetime without EOR of ~50 years [1]. The growing demand for petroleum is also being met in part by the increased reliance on production from proven unconventional oil reserves, for example, the Canadian oil sands, which has an estimated 300 billion barrels of ultimate potential recoverable reserve (heavy oil), the third largest reserve behind Venezuela and Saudi Arabia [2].

Successful EOR and oil sands operations rely on controlling the process fluid chemistry to favorably affect the mechanisms that govern oil droplet dewetting and liberation. In this paper, we will provide an overview of the scientific principles influencing oil droplet dynamics on solid surfaces, and extend the discussion to demonstrate how those governing mechanisms can be influenced by the commonly studied surface/interfacially active components namely surfactants and nanoparticles. The step-by-step process by which oil detaches from the solid surface can be summarized in sequence: (i) oil film thinning and rupture; (ii) oil de-wetting (recession) on the solid surface; and (iii) oil-solid surface adhesion and liberation.

Before describing the underlying principles that govern each step, it is worth considering the likely interaction between the oil and solid surface; i.e., the reservoir wettability. The reservoir environment can be characterized as either: (i) water-wet (water droplet contact angle, \( \theta = 0^\circ \) to \( 70^\circ \)); (ii) oil-wet (\( \theta \approx 110^\circ \) to \( 180^\circ \)); and (iii) neutrally-wet (\( \theta \approx 70^\circ \) to \( 110^\circ \)) exhibiting a similar affinity to both water and oil [3–5]. While it is understood that most reservoir environments were initially water-wet, the reservoir rock can evolve to become more oil-wet due to the deposition/adsorption of several indigenous organic polar species (asphaltenes, resins and naphthenic acids) present in crude oil [6–9]. For oil-wet reservoirs, oil recovery is poor due to no capillary imbibition. Hence, one of the criteria for successful EOR is to enhance capillary imbibition and reverse the wettability change by using chemical additives, although complete reversal to strongly water-wet surfaces is not favored for EOR [10]. An oil layer on a hydrophilic or hydrophobic solid surface is the basis for the following discussions.

2. Background Science

Oil recovery from the reservoir rock occurs by either displacement from squeezing or oil film thinning and rupture to form discrete oil droplets (Figure 1) that are removed by shear; the latter is of interest here.

![Figure 1. Schematic showing the four stages of oil film dewetting from a uniform thick film (I) to film thinning (II); formation of discrete oil patches (III); and recession of oil patches to form oil droplets at the new equilibrium wetting condition (IV).](image)

The long-time transformation from water-wet to an oil-wet reservoir occurs following the collapse of a thin aqueous layer separating the solid surface and oil layer. The stability of the thin water layer is attributed to the disjoining pressure that accounts for surface forces between the solid-water and water-oil interfaces. The total disjoining pressure (\( \Pi \)) includes contributions from electrostatic (\( \Pi_{el} \)), van der Waals (\( \Pi_{vdW} \)), and structural (\( \Pi_{st} \)) forces [11]

\[
\Pi = \Pi_{el} + \Pi_{vdW} + \Pi_{st}
\]

with the thin water layer collapsing when \( \Pi \) is negative. The disjoining pressure as a function of aqueous layer thickness has been calculated for a silica/water/oil (bitumen) system of salinity 1 mM KCl and pHS 3, 5, and 9, see Figure 2 (only \( \Pi_{el} \) and \( \Pi_{vdW} \) have been considered). While \( \Pi_{vdW} \) depends on the interaction Hamaker constant, \( \Pi_{el} \) is sensitive to pH and salinity, with the magnitude of the
electrostatic force dependent on the surface (zeta) potentials of silica and bitumen, and the Debye length. For crude oil, the pH dependent surface potentials result from ionization and surface activity of natural surfactants (naphthenic acids) [12–14]. At higher pHs, dissociation of the carboxylic-type surfactant increases the surface potential (negative) of the oil-water interface, with the magnitude increasing as more surfactant partitions at the interface. The high surface potentials at pH 9 form very stable thin-water layers, whereas in more acidic conditions, the disjoining pressure maxima decrease, and the thin-water layer in pH 3 is entirely unstable.

The stability of thick oil films is governed by the balance of gravity and capillary forces, with instability and the formation of discrete oil patches having been described analytically by Sharma [17], with the critical film thickness \( h_{cr} \) (Equation (2)) dependent on the oil-water interfacial tension \( \gamma_{o/w} \), the three-phase contact angle (\( \theta \)), and the minimum radius of the hole \( (r_m) \)

\[
h_{cr} = r_m \ln \left( \frac{2 \sin(\pi - \theta)}{\cos(\pi - \theta)} \right) \sqrt{\frac{\gamma_{o/w}}{2 \rho g}}
\]

where \( g \) is the acceleration due to gravity and \( \rho \) the density of oil. For a typical oil-wetted solid surface of three-phase contact angle of 145° and \( \gamma_{o/w} = 30 \) mN/m, \( h_{cr} \) is 0.05 mm for a stable minimum hole radius of 10 \( \mu \)m. Dependence on the fluid and surface properties is rather weak over the range of general applicability, with \( h_{cr} \) strongly influenced by the size of the stable hole in the oil film [16]. For very thick films \( (h \gg h_{cr}) \), thinning of the oil film is needed for dewetting, otherwise holes formed in the film will spontaneously collapse. The mechanisms for film thinning have not been extensively considered but are most likely to result from fluid shear in confined environments. Other factors that can influence the onset of film rupture include gas bubbles trapped in the oil film [18], and surface asperities that lead to non-uniform film thickness.

**Figure 2.** Disjoining pressure \( (\Pi = \Pi_{el} + \Pi_{vdW}) \) as a function of thin-water layer thickness \( (h) \) and pH. Zeta potentials \( (\psi) \) at pH 3, 5 and 9 are: 2.5, -55.6 and -78.2 mV for oil (bitumen) [15], and -12, -30 and -38 mV for silica, respectively. The Hamaker constant \( (\mathcal{A}_{SWB}) \) for silica/water/oil system is \( 5.7 \times 10^{-21} \) J [16]. \( \Pi_{vdW} = \frac{A_{vdW}}{6h^3}, \Pi_{el} = \frac{1}{2} \varepsilon_0 \varepsilon \kappa^2 \cos h(xh) - \frac{\kappa^2 - \kappa_0^2}{\sin^2 h(xh)} \), where \( \varepsilon \) and \( \varepsilon_0 \) are the dielectric permittivity of vacuum and relative dielectric permittivity of water, respectively. \( \kappa \) is the Debye length, which accounts for changes in salinity.
With the oil film ruptured, the circular hole begins to expand at a rate dependent on the fluid and interfacial properties (to be discussed below) [19]. Away from equilibrium, the process of droplet dewetting is driven by a change in energy following the creation of a new solid-water interface and the loss of oil-solid interface, assuming the change in oil-water interface during droplet recession can be considered negligible, that is

$$\frac{dG}{dA} = \gamma_{S/W} - \gamma_{O/S}$$  \hspace{1cm} (3)

where $\gamma$ is the interfacial tension and subscripts S, W, and O describe the solid, water and oil phases, respectively. Equation (3) can be simplified by the Young’s equation for an oil droplet on a solid surface given by

$$\cos \theta = \frac{\gamma_{O/W} - \gamma_{S/W}}{\gamma_{O/W}}$$ \hspace{1cm} (4)

to express the energy change during oil recession in terms of the equilibrium contact angle and oil-water interfacial tension (two measurable properties)

$$\frac{dG}{dA} = -\gamma_{O/W} \cos \theta.$$ \hspace{1cm} (5)

With $\gamma_{O/W}$ always greater than zero, Equation (5) confirms that oil recession is a spontaneous process when $\theta < 90^\circ$; i.e., the wetted solid surface is more water-wet (hydrophilic). The simple form of Equation (5) provides fundamental insight for effective EOR, highlighting the value of modifying surface wettability and oil-water interfacial tension. The smaller the $\theta$, the more favorable the condition for oil recession. Once the oil droplet has reached equilibrium, the work of adhesion ($W_A$) between oil and solid surface must be exceeded to liberate the oil droplet. By the reduction in area of oil-solid interface and generation of oil-water and solid-water interfaces, $W_A$ is given by

$$W_A = \gamma_{S/W} + \gamma_{O/W} - \gamma_{O/S}$$ \hspace{1cm} (6)

which when combined with the Young’s equation leads to

$$W_A = \gamma_{O/W}(1 - \cos \theta) \geq 0.$$ \hspace{1cm} (7)

With the unlikely condition of $\theta = 0$ for spontaneous liberation (droplet detachment from the solid surface), Equation (7) confirms the need for energy to detach oil droplets from the wetted surface. In order to detach an oil droplet from the solid surface the hydrodynamic lift force must exceed the contributions from the body and adhesion forces. An approximation of the adhesion force for a partially wetting droplet is, $F_A = \pi r \gamma_{O/W} \sin(\pi - \theta)$, where $r$ is the radius of oil-solid surface contact area [20]. The contour map in Figure 3 indicates the strongest adhesion (red color) when the oil-water interfacial tension and water droplet contact angle are high. Therefore, reducing both the oil-water interfacial tension and oil-water-solid three-phase contact angle leads to more favorable oil droplet liberation.
3. Dynamics of oil Film Recession

After creation of a hole on the oil film, the oil film recedes rapidly, governed by the receding force, \( F_R = \gamma_{O/W}[\cos(\pi - d) - \cos(\pi - \theta)], \) acting at the three-phase contact line \((\theta_d\) is the dynamic contact angle), with \( F_R \) diminishing towards the new equilibrium wetted-state, hence, the velocity of the three-phase contact line decreases with time. The dynamics of oil displacement on a solid surface are frequently described using the (i) hydrodynamic (HD); (ii) molecular-kinetic (MK); or (iii) combined models.

For more viscous fluids, such as crude oil, the hydrodynamic model relies on the solution of creeping flow in the vicinity of the three-phase contact line, with the no slip boundary condition relaxed to allow for finite slipping of the fluid/fluid contact line on a solid surface. Considering an effective slip length \((L_S)\), Cox presented a comprehensive hydrodynamic solution by segmenting the dynamic three-phase contact line into inner, intermediate, and outer regions, and correlated the apparent contact angle to the three-phase contact line displacement velocity, \( U [21,22] \)

\[
U = \frac{\gamma_{O/W}}{9 \mu_o} \left[ (\pi - \theta)^3 - (\pi - \theta_d)^3 \right] \left[ \ln \left( \frac{L}{L_S} \right) \right]^{-1}
\]

where \( \mu_o \) is the oil viscosity, \( \theta \) is the contact angle measured through the water phase, and \( L \) and \( L_S \) are the characteristic length of the oil droplet and the slip length, respectively. While determination of the slip length is nontrivial, the term is often used as a fitting parameter of the experimental data.

The molecular-kinetic model accounts for molecular displacements (adsorption/desorption) in the vicinity of the dynamic three-phase contact line. The model assumes that the solid surface behaves as a source of identical adsorption sites, and liquid molecules can detach and attach to neighboring sites by overcoming an energy barrier to molecular displacements [23]. The work to overcome the energy barrier is provided by a driving force governed by \( \gamma_{O/W} \) and an imbalance between the equilibrium and dynamic wetting states. The three-phase contact line displacement is described in terms of molecular displacement, defined as the distance between adsorption sites \( \lambda \) and a frequency \( (\kappa^0) \) of adsorption/desorption events at equilibrium, as shown in Figure 4.

![Figure 3. Apparent adhesion force for a partially wetting droplet (droplet volume = 10 μL).](image-url)
The relationship between the dynamic contact angle and the three-phase contact line velocity is given by

\[ U = 2 \lambda^0 \sin h \left( \frac{\gamma_{OW} \cos(\pi - \theta_d) - \cos(\pi - \theta)}{2k_B T} \right) \]  

(9)

where \( k_B \) is the Boltzmann constant and \( T \) the absolute temperature. The molecular displacement parameters (\( \lambda \) and \( \lambda^0 \)) are often combined and treated as the coefficient of contact-line friction, \( \zeta = \frac{k_B T}{\lambda \lambda^0} \), to describe the energy dissipated at the three-phase contact line, and neglecting any viscous dissipation in the bulk liquid \([24,25]\). Similar to the HD model, \( \zeta \) is treated as an adjustable parameter of the experimental data. Simplification of Equation (9) then follows when the \( \sin h \) function is small; i.e., not far from equilibrium—and Equation (9) reduces to the linear form

\[ U = \frac{\gamma_{OW}}{\zeta} \left[ \cos(\pi - \theta_d) - \cos(\pi - \theta) \right] \]  

(10)

Since each model neglects a contributing factor, a combined model approach can be considered to account for both the contact-line friction and viscous dissipation. As described by de Gennes and Brochard-Wyart \([26,27]\), the combined model for contact-line displacement is given by

\[ U = \frac{\gamma_{OW} \cos(\pi - \theta_d) - \cos(\pi - \theta)}{\zeta + \frac{3
u}{2
T} \ln \left( \frac{h}{\lambda^0} \right)} \]  

(11)

The sequence of images in Figure 5 show the dewetting process for an oil droplet deposited on a solid surface. In this example, a 10 µL droplet of extra heavy oil (13.6° API at 20 °C; SARA: 7.4% saturates, 37.8% aromatics, 15.3% resins, and 39.5% asphaltenes) was deposited on a glass substrate with a water contact angle <5°. Since the oil viscosity was ~6700 mPa-s at 20 °C, the substrate was heated to ~50 °C to promote faster spreading of the oil droplet on the solid surface. With the oil droplet at the equilibrium wetted-state, Milli-Q water was pumped underneath the oil-wetted solid surface at 1400 mL/min to completely submerge the oil droplet. The measurement cell temperature was maintained using a circulating water bath. Since \( \frac{dG}{dA} < 0 \), oil droplet recession occurs spontaneously and the oil-solid contact area reduced to attain a new equilibrium wetted-state, as described by Equation (4).
The rate of oil film dewetting can be determined from the dynamic contact angle, see Figure 6, with faster dewetting dynamics observed for higher temperature environments. Clearer differentiation between 60 °C and 80 °C is shown in the inset of Figure 6, with the new equilibrium wetted-states (oil-water-solid surface) attained within a few minutes, contrasting the 40 °C sample, which required more than 1 h to reach equilibrium. Moreover, the contact angles at equilibrium were shown to depend on temperature, decreasing from 63.7° to 54.1° and 51.3° with increasing temperature from 40 °C to 60 °C and 80 °C, respectively. Equation (4) shows that changes in the equilibrium wetted-state result from a change in the balance of energies acting on the three interfaces. Measuring \( \gamma_{O/W} \) at equivalent temperatures, Figure 7 confirms a small decrease in \( \gamma_{O/W} \) with increasing temperature. Hence, if it were assumed that \( \gamma_{O/S} \) and \( \gamma_{S/W} \) remained independent of temperature, then \( \theta \) would decrease in good agreement with Equation (4). Previous studies showed variation in the oil-water interfacial tension as a function of pH and temperature \([12,13,28–31]\), with the effect attributed to the partial solubility of naphthenic acids in water \([32,33]\).

![Figure 5](image-url)  
**Figure 5.** Time-dependent dewetting of an extra heavy oil droplet on a hydrophilic solid surface. The solid surface and water temperature were maintained at 40 °C. Images were captured at 20 fps for the first 15 min and 2 fps thereafter. Data captured using the Theta tensiometer (Biolin Scientific).

![Figure 6](image-url)  
**Figure 6.** Dewetting dynamics of an extra heavy oil film immersed in Milli-Q water at different temperatures: 40 °C, 60 °C, and 80 °C. Inset is an expanded region of the initial dewetting dynamics to differentiate between the two higher temperatures. Each experimental condition was repeated four times with measurement variability considered to be negligible.
Naphthenic acids are considered to be cyclic carboxylic acids of the general form R–COOH, where R can be any cyclo-aliphatic group [34]. Compared to asphaltenes, naphthenic acids are of lower molecular weight, typically less than 450 g/mol, spanning mainly C10 to C50 compounds with up to six fused ring structures that are mostly saturated [35]. Naphthenic acids preferentially adsorb on carbonate solid surfaces mainly by chemical interactions to modify the wettability from water-wet to oil-wet as the surface becomes saturated, although the process is reversible at elevated temperatures [36,37].

The initial receding rate (\(\frac{d\theta}{dt}\)) of the oil film was compared for each temperature with the rates correlated to changes in \(\gamma_{O/W}\) and \(\mu_o\), see Figure 7. Based on the HD model for contact line displacement (Equation (8)), which includes both parameters, the oil viscosity is the rate dependent parameter since the change in oil viscosity (−80.9%) with temperature is more significant than that of oil-water interfacial tension (−9.9%); between 60 °C and 80 °C the oil viscosity decreased by 38.9% and the initial receding rate increased by 107.2%. The same oil displacement data was fitted to both the HD and MK models (Figure 8). A least-squares difference between the experimental and theoretical \(\theta_d\) was made

\[
\Delta = \sum_{t=0}^{t_e} (\theta_{d,t,m} - \theta_{d,t,e})^2
\]

(12)

where \(\theta_{d,t,m}\) and \(\theta_{d,t,e}\) are the theoretical and experimental dynamic contact angles at time t, respectively, and the model fitting parameters were determined by minimizing the least-squares value.

During the process of oil film dewetting, the model fits appear in reasonable agreement with the experimental data. Slight variation is magnified at higher temperatures when the receding dynamics can be considered rapid for extra heavy crude oil, and experimental variability is more evident. The adjustable fitting parameters for each model (HD—\(\ln (L_c)\), MK—\(\zeta\)) reduced with increasing temperature, suggesting that the slip length of fluid/fluid contact line on a solid surface (\(L_c\)) increases and the coefficient of contact-line friction decreases when the oil viscosity is reduced, in good agreement with previous findings [38,39]. While our study only considered dewetting dynamics in Milli-Q water the effect of water chemistry on oil film dewetting has received little attention and is an area for

Figure 7. Initial oil droplet receding rate as a function of temperature, correlated to changes in oil viscosity and oil-water interfacial tension (error ± 0.03 mN/m). Symbols: circle—oil droplet receding rate, triangle—\(\gamma_{O/W}\), square—\(\mu_o\).
further study. High salinity brine can increase oil-water interfacial tension \[\gamma_{O/W}\] and enhance solvation forces between solid-liquid interfaces \[\gamma_{S/L}\], bind surfactants to substrate via multicomponent ion exchange \[\gamma_{S/L}\], and impact the stability of chemical additives used for EOR.

![Figure 8](image)

**Figure 8.** Hydrodynamic (HD) and molecular-kinetic (MK) model fittings of oil film dewetting at 40 °C (a), 60 °C and 80 °C (b). Shaded lines represent the experimental data and the HD and MK models identified by the solid and dash lines, respectively. (c) Optimal fitting parameters, HD (closed symbols)—ln \((\frac{L}{L_0})\), MK (open symbols)—\(\zeta\).

### 4. Surfactant Oil Droplet Displacement

Surfactants are widely used in EOR to reduce \(\gamma_{O/W}\) and enhance water-wetting of the solid surface. Surfactants are often described as amphiphilic molecules composed of a hydrophilic head and a hydrophobic tail, thus surfactants favorably partition at solid-liquid and liquid-liquid interfaces. The accumulation of surfactants at an interface is a function of the surfactant concentration in the bulk fluid as described by the general form of the Gibbs’ adsorption equation for a binary, isothermal system, 

\[
d\gamma = -RT\Gamma C_s \ln \Gamma_s
\]

where \(\Gamma_s\) is the surface excess of surfactant, \(C_s\) the surfactant concentration in the bulk fluid, and \(RT\) the thermal energy of the system. As a function of concentration, surfactants in solution exist in the monomer-form at low surfactant concentrations, reaching a concentration of maximum solubility of the monomer-form, forming micelles via self-association. This concentration is termed the critical micelle concentration (CMC). Surfactant adsorption and displacement of organic species on solid surfaces and the resultant wettability modification is dependent on the surfactant concentration. At extremely low concentrations, surfactant monomers adsorb as individual molecules with no interaction between the adsorbed molecules. At higher concentrations (<CMC) surfactant molecules associate to form patchy hemi-micelles on the solid surface, with surfactants coordinating in the tail-tail confirmation. Further increases in concentration lead to saturation of all available surface sites and the formation of a surfactant bi-layer at the CMC \[\gamma_{S/L}\]. Formation of a bi-layer would orientate the surfactant hydrophilic head group away from the solid surface, thus increasing the water-wetting nature of the reservoir rock, favorable for oil droplet displacement (Equations (5) and (7)). Mechanisms for wettability modification by different surfactants are described below.

Composition of the reservoir surface (sandstone, carbonate, and deposited organic species) often dictates the surfactant selection for wettability modification, with surfactants categorized as cationic, anionic and non-ionic, based on the charge characteristics of hydrophilic groups. Surfactant adsorption on the solid surface can occur via electrostatic and van der Waals forces, and hydrogen bonding, with the extent of wettability modification a function of several properties including surfactant adsorption kinetics, surfactant structure, temperature, pH, salinity. A brief summary of surfactants considered...
for EOR is provided in Table A1, with remarks provided for changes in solid surface wettability and interfacial tension.

While electrostatic interactions are often considered to describe surfactant-solid surface adsorption, such simplicity does not describe the potential for surfactants to modify solid surface wettability, when many other factors such as oil saturation, clay content, divalent cations, pH, and temperature influence the action of the surfactant.

Cationic surfactants are frequently used to treat carbonate reservoirs and include permanently charged ammonium groups (ammonium bromide and ammonium chloride) [44]. Adsorbed polar components of crude oil (i.e., negatively charged naphthenic acids) can be removed from the solid surface by forming ion pairs with cationic surfactants via strong ionic interaction. Removal of contaminants transforms the solid surface wettability to more water-wet [44,45]. The use of cationic surfactants to treat sandstone has also been demonstrated, although the chemical effectiveness in carbonate reservoirs is greater [46].

Anionic surfactants including sulfates, sulfonates, phosphates, and carboxylates, have been shown to modify wettability in both carbonate and sandstone reservoirs. Wettability modification occurs via two mechanisms [44,45]: (i) anionic surfactants interact with the organic species via hydrophobic forces, exposing the surfactant head group to make the solid surface more water-wet (wettability modification for sandstone reservoirs); and (ii) via strong electrostatic forces with carbonate surfaces, anionic surfactants can displace organic species exposing the underlying water-wet surface [46].

Non-ionic surfactants such as alcohols, esters and ethers have been used to modify the wettability of carbonate and sandstone surfaces [47], being highly effective in high salinity water. With no contribution from electrostatic forces, non-ionic surfactants interact via hydrophobic forces with deposited organic species, and hydrogen bonding with hydroxyl groups on the solid surface [48,49]. Research has shown that non-ionic surfactants can modify highly oil-wet carbonate to weakly oil-wet or even water-wet ($\theta < 80^\circ$) following the addition of 0.1 wt % surfactant [50].

Sodium dodecyl sulfate (SDS), an anionic surfactant, was used to displace an oil film deposited on a glass substrate at 60 °C (Figure 9a). Adding SDS to the aqueous phase reduced $\gamma_{O/W}$, and the CMC was measured at ~0.1 wt % (3.5 mM) at 60 °C (Figure 9c). Increasing the SDS concentration from $5 \times 10^{-4}$ wt % to $5 \times 10^{-3}$ wt % increased both the rate of oil film displacement and equilibrium oil droplet contact angle (inferred from lower water contact angle, $\theta$). The equilibrium contact angle reduced from 54.1° in the absence of SDS to 48.7° and 36.5° for $5 \times 10^{-4}$ wt % and $5 \times 10^{-3}$ wt % SDS, respectively. Fitting the HD and MK models confirmed an increased slip length ($L_S$) and reduced coefficient of contact-line friction ($\zeta$) at higher SDS concentrations. Figure 9b illustrates the benefit of injecting surfactants at a concentration greater than the CMC. The very low oil-water interfacial tension (~5.75 mN/m) causes the oil film to continually recede and eventually detach from the solid surface when the oil droplet buoyant force (3.77 µN) exceeds the solid surface-oil droplet adhesion force (2.01 µN).
5. Nanoparticle Oil Droplet Displacement

The application of ultra-small particles (nanoparticles) to enhance oil film displacement has been demonstrated. Nanoparticles are typically 1 to 100 nm and are ideal for EOR applications with particle sizes smaller than the pore diameter, hence nanofluids flow through the porous media without obstructing the porous network. In addition, their high surface area to volume ratio increases their effectiveness at low particle concentrations, and promotes their kinetic stability [51]. An overview of nanoparticles (nanofluids) used to displace oil films is provided in Table A2.

For oil film displacement, different nanoparticles have been considered including metal oxides, organic, inorganic, and composite particles. Metal oxides nanoparticles (Al₂O₃, CuO, TiO₂ and Fe₂O₃) have been shown to lower γₒ/W [52] and increase the disjoining pressure between the solid surface and oil-water interface [53]. The interfacial tension decreases as nanoparticles partition at the oil-water interface from the aqueous phase. Unlike surfactants, nanoparticles are not amphiphilic and their affinity to partition at an oil-water interface is governed by their particle size and surface wettability. The general expression of \( E = \pi a^2 \gamma \theta (1 \pm \cos \theta)^2 \), describes the particle detachment energy from an oil-water interface (describes detachment into either liquid phase), where \( a \) is the particle radius [54]. When \( \cos \theta = 0 \) the particle detachment energy is maximized, although for nanoparticles \( E \) can be of the order of a few \( k_BT \) which can result in reversible adsorption, similar to a surfactant molecule.

The reduction in oil-water interfacial tension has been shown to correlate to the nanoparticle size and the particle specific surface area [55]. Al₂O₃ nanoparticles of diameter 20 nm and 45 nm were shown to lower the oil-water interfacial tension to ~13.6 mN/m and ~8.6 mN/m, respectively [56]. Compared to surfactants, interfacial tension reduction by nanoparticles is often smaller. For example, silica nanoparticles (7–14 nm) dosed at 0.01–0.10 wt % reduced \( \gamma_{O/W} \) to ~10 mN/m from ~15–20 mN/m [57–59], while TiO₂ nanoparticles (58 nm) reduced \( \gamma_{O/W} \) from 23 mN/m to 18 mN/m when dosed at 0.01–0.05 wt % [53]. As such, enhanced oil film displacement by nanoparticles is likely to occur via other mechanisms; i.e., structural disjoining pressure and wettability modification.

**Structural disjoining pressure** is a consequence of nanofluids exhibiting super-spreading behavior. Nanoparticles self-assemble in the vicinity of the three-phase contact line to form a liquid wedge at the de-pinning point, see Figure 10. As nanoparticles accumulate in the liquid wedge a structural disjoining
pressure (Equation (14)) gradient is established with the highest pressure at the oil droplet-solid surface vertex, driving the nanofluid to spread and cause the oil film to recede. As explained by Wasan and co-workers [60,61], the spreading coefficient ($S$) of the nanofluid is determined by the sum of the capillary pressures at the equilibrium film thickness ($\Pi_0(h_e)$) and disjoining pressure ($\Pi(h)$)

$$S = \Pi_0(h_e)h_e + \int_{h_e}^{\infty} \Pi(h) dh$$

(13)

![Figure 10](image_url)

Figure 10. Schematic showing the ordered accumulation of nanoparticles to form a liquid wedge. The structural disjoining pressure increases towards the de-pinning point. The structural disjoining pressure exceeds the Laplace pressure, deforming the meniscus profile as represented by inner and outer contact lines. The contribution from the long range structural disjoining pressure dominates the short range electrostatic and van der Waals forces. The figure has been modified from Zhang et al. [62] and Chengara et al. [11].

When the thickness of the liquid wedge exceeds one particle diameter, nanoparticles accumulate in ordered layers. This layered arrangement of nanoparticles increases the excess pressure in the liquid wedge with the structural disjoining pressure described based on the theory of thin liquid films [63]:

$$\Pi_{st}(h) = \Pi_0 \cos(\omega h + \varnothing_2) e^{-\kappa_1 h} + \Pi_1 e^{-\delta(h-d)}$$

(14)

$$\Pi_{st}(h) = -P, \ 0 < h < d$$

where $d$ is the nanoparticle diameter, $P$ the osmotic pressure of nanofluid, and all other parameters ($\Pi_0$, $\Pi_1$, $\omega$, $\varnothing_2$, $\kappa$ and $\delta$) are fitted as cubic polynomials varying with particle concentration. Contributions from van der Waals, electrostatic and structural forces have been considered by Chengara et al. [11] (Figure 10). The structural forces are long range and govern the behavior of thick liquid films, with nanoparticle size, concentration, temperature, and fluid salinity, all contributing to the magnitude of the structural disjoining pressure.

Wettability modification by nanoparticles enhances oil droplet displacement when nanoparticles deposit on the solid surface. The deposition/adsorption is influenced by electrostatic forces, with the nanoparticle decorated solid surface more water-wet due to deposition of hydrophilic particles to form a
heterogeneous surface and increased nano/micron-scale roughness [64–68]. Wettability of heterogeneous surfaces has been described by Cassie-Baxter [69], with the apparent contact angle on a composite solid surface given by, \[ \cos \theta_{CB} = f_1 \cos \theta_1 + f_2 \cos \theta_2, \] where \( f_1 \) is the fractional area of the surface with contact angle \( \theta_1 \), \( f_2 \) is the fractional area of the surface with contact angle \( \theta_2 \), and \( \theta_{CB} \) is the Cassie-Baxter contact angle. The Cassie-Baxter model can be combined with the Wenzel wetting model [70] to account for surface roughness effects, \[ \cos \theta_{W} = R' \cos \theta_{CB}, \] where \( R' \) is the ratio of the true area of the solid to its planar projection. With \( R' \) always greater than 1, the Wenzel model confirms nano/micron-scale roughness lowers the contact angle of a water-wet surface, thus increasing the potential for oil droplet displacement. For example, metal oxide nanoparticles (ZrO\(_2\) and NiO < 50 nm) were shown to deposit on an oil-wet surface modifying the contact angle from 152° (untreated surface) to 44° and 86° for ZrO\(_2\) and NiO, respectively. The mean roughness of those surfaces was shown to increase from 70.6 nm (untreated surface) to 2.32 \( \mu \)m (ZrO\(_2\) treated surface) and 330 nm (NiO treated surface) [71].

Oil film displacement can be enhanced when nanoparticles are mixed with surfactants. Fluid blends lower the oil-water interfacial tension below a surfactant only system, with surfactants increasing the interfacial activity of nanoparticles [72]. The decrease in oil-water interfacial tension depends on the surfactant-particle interaction and surfactant concentration [73]. The effect of nanoparticles is lessened at surfactant concentrations greater than the CMC. The use of surfactant blends and composite particles (polymer-coated particles) to enhance oil film displacement have also been considered but such mechanisms are considered outside the scope of this paper [74,75]. Recent studies, which have considered composite fluids (particles), have been summarized in Table A3.

6. Conclusions

While demand for oil continues to rise, challenges in extraction become ever more complex. Extraction from confined, unfavorable environments, and production of unconventional oil is increasing the dependence on alternative extraction methods to deliver enhanced oil recovery. Often the interaction between the oil and solid surface limits recovery with oil strongly adhered to an oil-wet surface. Oil film recession is spontaneous when the solid surface is water-wet, and the adhesion force to be overcome to liberate an oil droplet from a solid surface diminishes with decreasing contact angle and oil-water interfacial tension. The rate of oil film recession and oil droplet equilibrium contact angle can be modified through the careful selection of chemicals. Surfactants have extensively been considered and used in production to lower oil-water interfacial tension and modify the solid surface to more water-wet. An alternative mechanism for oil film displacement has been identified when using nano-fluids. Accumulation of nanoparticles in a liquid wedge between oil and solid surface results in a long range structural disjoining pressure gradient causing the three-phase contact line to move (i.e., oil film recede due to super-spreading of the nano-fluid).

Controlling interfacial behavior in the reservoir provides a route for enhanced oil recovery. Significant research effort is ongoing to design more effective chemicals that perform in challenging environments (temperature, pressure, salinity, clays), deliver performance at the targeted site (i.e., minimize material loss), and do not impact the environment. Enhanced oil recovery will ensure effective utilization of crude oil resources, and the fundamental mechanisms governing oil film displacement and oil droplet detachment are underpinned by knowledge of interfacial and colloidal forces.

Author Contributions: S.T. performed the experiments; S.T. and D.H. analyzed the data; S.T. and D.H. drafted the manuscript; T.V.J.C. and D.P. reviewed the final manuscript.

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Conflicts of Interest: The authors declare no conflict of interest.
## Appendix

<table>
<thead>
<tr>
<th>Surfactants</th>
<th>Conc.</th>
<th>Solid Surface</th>
<th>Oil Type</th>
<th>Remarks</th>
<th>Ref.</th>
</tr>
</thead>
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<tr>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>(n)-C\textsubscript{8}N(CH\textsubscript{3})\textsubscript{3}Br (C8TAB) in brine</td>
<td>4.0 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 57(^\circ), IFT = 2.85 mN/m</td>
<td>[44]</td>
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<tr>
<td>(n)-C\textsubscript{10}N(CH\textsubscript{3})\textsubscript{3}Br (C10TAB) in water</td>
<td>0.4 wt %</td>
<td>Calcite</td>
<td>Decane mixed with naphthenic acids</td>
<td>IFT = 2.67 mN/m</td>
<td>[45]</td>
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<tr>
<td>(n)-C\textsubscript{12}N(CH\textsubscript{3})\textsubscript{3}Br (C12TAB) in water</td>
<td>0.4 wt %</td>
<td>Calcite</td>
<td>Decane mixed with naphthenic acids</td>
<td>IFT = 0.59 mN/m</td>
<td>[45]</td>
</tr>
<tr>
<td>(n)-C\textsubscript{12}N(CH\textsubscript{3})\textsubscript{3}Br (C12TAB) in brine</td>
<td>5.0 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 12(^\circ), IFT = 0.81 mN/m</td>
<td>[44]</td>
</tr>
<tr>
<td>(n)-C\textsubscript{12}N(CH\textsubscript{3})\textsubscript{3}Br (C12TAB) in brine</td>
<td>1.0 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 27(^\circ), IFT = 0.38 mN/m</td>
<td>[44]</td>
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<tr>
<td>(n)-Decyl triphenylphosphonium bromide (C10TPPB) in water</td>
<td>0.4 wt %</td>
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<td>Decane mixed with naphthenic acids</td>
<td>IFT = 3.56 mN/m</td>
<td>[45]</td>
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<tr>
<td>Cetyltrimethylammonium bromide (CTAB) in brine</td>
<td>75–2620 ppm (0.0075–0.262 wt %)</td>
<td>Dolomite</td>
<td>Crude oil</td>
<td>Contact angle = 69(^\circ), IFT = 4.8 mN/m</td>
<td>[77]</td>
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<tr>
<td>Dodecyltrimethylammonium bromide (DTAB) in brine</td>
<td>0.06 wt %</td>
<td>Quartz</td>
<td>Crude oil</td>
<td>Contact angle = 95(^\circ), IFT = 2.49 mN/m</td>
<td>[78]</td>
</tr>
<tr>
<td>(n)-(C\textsubscript{8}-C\textsubscript{18})-N(CH\textsubscript{3})\textsubscript{2}(CH\textsubscript{2}-Ph)Cl (ADMBACl) in brine</td>
<td>0.5 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 26(^\circ), IFT = 0.41 mN/m</td>
<td>[44]</td>
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<tr>
<td>(n)-C\textsubscript{8}Ph-(EO)\textsubscript{2}-N(CH\textsubscript{3})\textsubscript{2}(CH\textsubscript{2}-Ph)Cl (Hyamine) in brine</td>
<td>0.2 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 21(^\circ), IFT = 0.48 mN/m</td>
<td>[44]</td>
</tr>
<tr>
<td>Coconut oil alkyl trimethylammonium chloride (ARQUAD C-50) in water</td>
<td>0.4 wt %</td>
<td>Calcite</td>
<td>Decane mixed with naphthenic acids</td>
<td>IFT = 0.53 mN/m</td>
<td>[45]</td>
</tr>
<tr>
<td>Trimethyl tallowalky ammonium choride (ARQUAD T-50) in water</td>
<td>0.4 wt %</td>
<td>Calcite</td>
<td>Decane mixed with naphthenic acids</td>
<td>IFT = 0.69 mN/m</td>
<td>[45]</td>
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<tr>
<td>Methyladecylbis ammonium tribromide</td>
<td>0.001–1 mM</td>
<td>Mica</td>
<td>Kerosene mixed with (n)-decane</td>
<td>Contact angle = 87(^\circ), IFT = 0.18 mN/m</td>
<td>[79]</td>
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Table A1. Cont.

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<th>Surfactants</th>
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<th>Oil Type</th>
<th>Remarks</th>
<th>Ref.</th>
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<td>Anionic surfactants</td>
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<td>$n$-((C_{12}-C_{15})-(EO)_{15})-SO$_3$Na (S-150) in brine</td>
<td>0.5 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 63$^\circ$, IFT = 2.29 mN/m</td>
<td>[44]</td>
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<tr>
<td>$n$-C$<em>{13}$-(EO)$</em>{8}$-SO$_3$Na (B 1317) in brine</td>
<td>0.5 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 40$^\circ$, IFT = 0.78 mN/m</td>
<td>[44]</td>
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<tr>
<td>$n$-C$<em>8$-(EO)$</em>{2}$-SO$_3$Na (S-74) in brine</td>
<td>0.5 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 49$^\circ$, IFT = 6.72 mN/m</td>
<td>[44]</td>
</tr>
<tr>
<td>$n$-(C$<em>{12}$-C$</em>{15}$)-(PO)$<em>{4}$-(EO)$</em>{2}$-OSO$_3$Na (APES) in brine</td>
<td>1.0 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 44$^\circ$, IFT = 0.082 mN/m</td>
<td>[44]</td>
</tr>
<tr>
<td>$n$-(C$_8$O$_2$CCH$_2$)((n)-C$_8$O$_2$C)CH-SO$_3$Na (Cropol) in brine</td>
<td>0.5 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 55$^\circ$, IFT = 8.77 mN/m</td>
<td>[44]</td>
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<tr>
<td>Sodium dodecyl sulfate (SDS) in brine</td>
<td>0.1 wt %</td>
<td>Chalk</td>
<td>Crude oil mixed with heptane</td>
<td>Contact angle = 39$^\circ$, IFT = 2.95 mN/m</td>
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<tr>
<td>Sodium dodecyl sulfate (SDS) in water</td>
<td>0.4 wt %</td>
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<td>Decane mixed with naphthenic acids</td>
<td>IFT = 4.77 mN/m</td>
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<tr>
<td>Sodium dodecyl 3EO sulfate in brine</td>
<td>0.05 wt %</td>
<td>Calcite</td>
<td>Crude oil</td>
<td>Contact angle ~45$^\circ$, IFT = 0.003 mN/m</td>
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<tr>
<td>Alkylphenol disulfonate in Na$_2$CO$_3$/NaCl</td>
<td>0.05 wt %</td>
<td>Calcite</td>
<td>Crude oil</td>
<td>Contact angle ~110$^\circ$, IFT = 0.0011 mN/m</td>
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<tr>
<td>Sodium nonyl phenol ethoxylated sulfate (4EO) in Na$_2$CO$_3$/NaCl</td>
<td>0.30 wt %</td>
<td>Calcite</td>
<td>Crude oil</td>
<td>Contact angle ~80$^\circ$, IFT = 0.00812 mN/m</td>
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<tr>
<td>Sodium dodecyl sulfate in Na$_2$CO$_3$/NaCl</td>
<td>0.05 wt %</td>
<td>Calcite</td>
<td>Crude oil</td>
<td>Contact angle ~60$^\circ$, IFT = 0.03 mN/m</td>
<td>[50]</td>
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<td>Polyether sulfonate in Na$_2$CO$_3$/NaCl</td>
<td></td>
<td></td>
<td>Crude oil</td>
<td>Contact angle ~40$^\circ$, IFT = 0.0001 mN/m</td>
<td>[50]</td>
</tr>
<tr>
<td>C$<em>{12}$-C$</em>{13}$ propoxy sulfate (8PO) in Na$_2$CO$_3$/NaCl</td>
<td>0.05 wt %</td>
<td>Calcite</td>
<td>Crude oil</td>
<td>Contact angle ~70$^\circ$, IFT = 0.116 mN/m</td>
<td>[50]</td>
</tr>
<tr>
<td>Alkylphenol disulfonate + C$_{14}$-T-isofol propoxy sulfonate (8PO) in Na$_2$CO$_3$/NaCl</td>
<td>0.075 wt %</td>
<td>Calcite</td>
<td>Crude oil</td>
<td>Contact angle ~70$^\circ$, IFT = 0.116 mN/m</td>
<td>[50]</td>
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<tr>
<td>Methyl alcohol+Proprietary sulfonate in brine</td>
<td>0.02–0.20 wt %</td>
<td>Shale (siliceous)</td>
<td>Crude oil</td>
<td>Contact angle ~38$^\circ$, IFT = 0.4 mN/m</td>
<td>[81]</td>
</tr>
<tr>
<td>Sodium laureth sulfate in brine</td>
<td>0.02–0.05 wt %</td>
<td>Quartz</td>
<td>Crude oil</td>
<td>Contact angle ~110$^\circ$, IFT = 2.007 mN/m</td>
<td>[76]</td>
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<tr>
<td>Sodium lauryl monoether sulfate in brine</td>
<td>0.035 wt %</td>
<td>Quartz</td>
<td>Crude oil</td>
<td>Contact angle ~116.1$^\circ$, IFT = 2.49 mN/m</td>
<td>[78]</td>
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Table A1. Cont.

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<th>Surfactants</th>
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<th>Remarks</th>
<th>Ref.</th>
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<td>Non-ionic surfactants</td>
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<td></td>
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<tr>
<td>Poly-oxyethylene alcohol (POA) in brine</td>
<td>750–1050 ppm (0.075–0.105 wt %)</td>
<td>Dolomite</td>
<td>Crude oil</td>
<td>IFT = 2.0 mN/m</td>
<td>[47]</td>
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<tr>
<td>Ethoxylated C11–C15 secondary alcohol (Tergitol 15-S-3) in water</td>
<td>0.4 wt %</td>
<td>Calcite</td>
<td>Decane mixed with naphthenic acids</td>
<td>IFT = 4.44 mN/m</td>
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<tr>
<td>Ethoxylated C11–C15 secondary alcohol (Tergitol 15-S-7) in water</td>
<td>0.4 wt %</td>
<td>Calcite</td>
<td>Decane mixed with naphthenic acids</td>
<td>IFT = 1.39 mN/m</td>
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<tr>
<td>Ethoxylated C11–C15 secondary alcohol (Tergitol 15-S-40) in water</td>
<td>0.4 wt %</td>
<td>Calcite</td>
<td>Decane mixed with naphthenic acids</td>
<td>IFT = 11.5 mN/m</td>
<td>[45]</td>
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<td>Nonylphenoxypoly(ethyleneoxy)ethanol (Igepal CO-530) in water</td>
<td>0.4 wt %</td>
<td>Calcite</td>
<td>Decane mixed with naphthenic acids</td>
<td>IFT = 0.33 mN/m</td>
<td>[45]</td>
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<td>C12–C15 linear primary alcohol ethoxylate (Neodol 25-7) in water</td>
<td>0.4 wt %</td>
<td>Calcite</td>
<td>Decane mixed with naphthenic acids</td>
<td>IFT = 2.02 mN/m</td>
<td>[45]</td>
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<tr>
<td>Secondary alcohol ethoxylate in Na2CO3/NaCl</td>
<td>0.10 wt %</td>
<td>Calcite</td>
<td>Crude oil</td>
<td>Contact angle ~20°, IFT = 0.0017 mN/m</td>
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<td>Nonyl phenol ethoxylate in Na2CO3/NaCl</td>
<td>0.10 wt %</td>
<td>Calcite</td>
<td>Crude oil</td>
<td>Contact angle ~80°, IFT = 0.0006 mN/m</td>
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<tr>
<td>Branched alcohol oxyalkylate in brine</td>
<td>0.02–0.20 wt %</td>
<td>Shale (siliceous)</td>
<td>Crude oil</td>
<td>Contact angle = 60°, IFT = 9.8 mN/m</td>
<td>[81]</td>
</tr>
<tr>
<td>Polyoxyethylene octyl phenyl ether in brine</td>
<td>0.04 wt %</td>
<td>Quartz</td>
<td>Crude oil</td>
<td>Contact angle = 95°, IFT = 4.05 mN/m</td>
<td>[76]</td>
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<td>Alkylpolyglycosides in brine</td>
<td>0.05 wt %</td>
<td>Quartz</td>
<td>Crude oil</td>
<td>Contact angle = 58.8°, IFT = 2.49 mN/m</td>
<td>[78]</td>
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</table>

*a* not all studies reported contact angle or interfacial tension data. 
b IFT is interfacial tension.
Table A2. Nanoparticles/fluids.

<table>
<thead>
<tr>
<th>Nanoparticles/Fluids</th>
<th>Solid Surface</th>
<th>Oil Type</th>
<th>Remarks (^a)</th>
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<td><strong>Metal oxides</strong></td>
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<tr>
<td>TiO₂ (0.01–1 wt %)</td>
<td>Sandstone</td>
<td>Heavy oil</td>
<td>Contact angle = 90°</td>
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<td>TiO₂ (0.01–0.10 wt %)</td>
<td>Sandstone</td>
<td>Heavy crude oil</td>
<td>Slight IFT reduction ~(\Delta \gamma = 1) mN/m</td>
<td>[52]</td>
</tr>
<tr>
<td>TiO₂ (0.01–0.05 wt %)</td>
<td>Sandstone</td>
<td>Heavy oil</td>
<td>Contact angle change from 127° to 81°, Slight IFT reduction</td>
<td>[53]</td>
</tr>
<tr>
<td>Al₂O₃ (0.01–0.10 wt %)</td>
<td>Sandstone</td>
<td>Heavy crude oil</td>
<td>Slight IFT reduction ~(\Delta \gamma = 1) mN/m</td>
<td>[52]</td>
</tr>
<tr>
<td>NiO (0.01–0.10 wt %)</td>
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<td>Heavy crude oil</td>
<td>Slight IFT reduction ~(\Delta \gamma = 1) mN/m</td>
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</tr>
<tr>
<td><strong>Organic</strong></td>
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<tr>
<td>Janus nanoparticles</td>
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<td>Hexane</td>
<td>IFT = 12 mN/m</td>
<td>[83]</td>
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<tr>
<td>(0.0025–0.0004 mM)</td>
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<tr>
<td>Carbon nanotubes</td>
<td>Glass</td>
<td>Crude oil</td>
<td>IFT reduction ~3 mN/m</td>
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<tr>
<td>(0.05–0.50 wt %)</td>
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<tr>
<td>Nanocellulose</td>
<td>Glass</td>
<td>Crude oil</td>
<td>IFT = 0.7 mN/m</td>
<td>[85]</td>
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<tr>
<td>(0.2–1.0 wt %)</td>
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<td></td>
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<tr>
<td><strong>Inorganic</strong></td>
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<tr>
<td>SiO₂ (0.1–0.6 wt %)</td>
<td>Carbonate</td>
<td>Crude oil</td>
<td>Contact angle = 51°</td>
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<td>SiO₂ (0.5–4.0 wt %)</td>
<td>Calcite (oil-wet)</td>
<td>n-decane</td>
<td>Contact angle = 20°</td>
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<tr>
<td>SiO₂ (0.1–5 wt %)</td>
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<td>Crude oil</td>
<td>Contact angle = 0°</td>
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<td>SiO₂ (0.025–0.2 wt %)</td>
<td>Calcite (oil-wet)</td>
<td>n-heptane</td>
<td>Contact angle = 41.7°</td>
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<td>SiO₂ (0.4 effective volume fraction)</td>
<td>Glass</td>
<td>Model oil</td>
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<tr>
<td>SiO₂ (0.01–0.10 wt %)</td>
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<td>Crude oil</td>
<td>Contact angle = 22°, IFT = 7.9 mN/m</td>
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<td>SiO₂ (0.10 wt %)</td>
<td>Sandstone</td>
<td>Light crude oil</td>
<td>Contact angle change from 34° to 32°, IFT reduced from 20 to 10 mN/m</td>
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\(^a\) Reference
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<tr>
<th>Nanoparticles/Fluids</th>
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<th>Oil Type</th>
<th>Remarks</th>
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<td>SiO$_2$ (0.01–0.10 wt %)</td>
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<td>Slight IFT reduction $\Delta \gamma = 1$ mN/m</td>
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</tr>
<tr>
<td>Hydrophilic silica (0.01–0.10 wt %)</td>
<td>Glass/Sandstone</td>
<td>Light crude oil</td>
<td>Contact angle $\sim 20^\circ$, IFT $\sim 8$ mN/m</td>
<td>[59]</td>
</tr>
<tr>
<td>Hydrophilic, neutralized, and hydrophobic silica (0.2–0.3 wt %)</td>
<td>Sandstone</td>
<td>Crude oil</td>
<td>Contact angle $\sim 35^\circ$</td>
<td>[57]</td>
</tr>
<tr>
<td>Hydrophobic silica (0.1–0.4 wt %)</td>
<td>Sandstone</td>
<td>Crude oil</td>
<td>Contact angle = 95.4°, IFT = 1.75 mN/m</td>
<td>[90]</td>
</tr>
<tr>
<td>Nanostructure particles (0.05–0.50 wt %)</td>
<td>Sandstone</td>
<td>Light crude oil</td>
<td>Wettability index = 0.36 (wettability index = 1 is water-wet)</td>
<td>[91]</td>
</tr>
<tr>
<td>Silica colloidal nanoparticles (0.05–0.50 wt %)</td>
<td>Sandstone</td>
<td>Light crude oil</td>
<td>Wettability index = 0.57 (wettability index = 1 is water-wet)</td>
<td>[91]</td>
</tr>
</tbody>
</table>

* not all studies reported contact angle or interfacial tension data. $^b$ IFT is interfacial tension. $^c$ NA is not available.
### Table A3. Composite fluids.

<table>
<thead>
<tr>
<th>Composite Fluids</th>
<th>Solid Surface</th>
<th>Oil Type</th>
<th>Remarks $^a$</th>
<th>Ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Blend systems</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>SDS and SiO$_2$ (Patented nanofluid—No reported concentration)</td>
<td>Glass</td>
<td>Crude oil</td>
<td>Contact angle = 1.2°</td>
<td>[62]</td>
</tr>
<tr>
<td>SDS and hydrophilic and hydrophobic SiO$_2$ (Surfactant: 100–6000 ppm, particle: 1000–2000 ppm)</td>
<td>Sandstone</td>
<td>Kerosene</td>
<td>IFT $^b$ = 1.81 mN/m</td>
<td>[72]</td>
</tr>
<tr>
<td>SDS and ZrO$_2$ (Surfactant: 0.001–5 CMC, particle: 0.001–0.050 wt %)</td>
<td>NA $^c$</td>
<td>$n$-heptane</td>
<td>IFT = 10 mN/m</td>
<td>[92]</td>
</tr>
<tr>
<td><strong>Composite nanoparticles</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zwitterionic polymer and SiO$_2$ (coated) (No reported concentration)</td>
<td>Sandstone</td>
<td>$n$-decane</td>
<td>IFT = 35 mN/m</td>
<td>[74]</td>
</tr>
</tbody>
</table>

$^a$ not all studies reported contact angle or interfacial tension data. $^b$ IFT is interfacial tension. $^c$ NA is not available.
References

4. Chilingar, G.V.; Yen, T.F. Some Notes on Wettability and Relative Permeabilities of Carbonate Reservoir Rocks, II. Energy Sources 1983, 7, 67–75. [CrossRef]
17. Sharma, A. Disintegration of macroscopic fluid sheets on substrates—A singular perturbation approach. J. Colloid Interface Sci. 1993, 156, 96–103. [CrossRef]

28. Schramm, L.L.; Smith, R.G. The influence of natural surfactants on interfacial charges in the hot-water process for recovering bitumen from the athabasca oil sands. *Colloids Surfaces* 1985, 14, 67–85. [CrossRef]

29. Drelich, J.; Miller, J.D. Surface and interfacial tension of the Whiterocks bitumen and its relationship to bitumen release from tar sands during hot water processing. *Fuel* 1994, 73, 1504–1510. [CrossRef]

30. Drelich, J.; Bukka, K.; Miller, J.D.; Hanson, F.V. Surface Tension of Toluene-Extracted Bitumens from Utah Oil Sands as Determined by Wilhelmy Plate and Contact Angle Techniques. *Energy Fuels* 1994, 8, 700–704. [CrossRef]


32. Rogers, V.V.; Liber, K.; MacKinnon, M.D. Isolation and characterization of naphthenic acids from Athabasca oil sands tailings pond water. *Chemosphere* 2002, 48, 519–527. [CrossRef]


45. Wu, Y.; Shuler, P.J.; Blanco, M.; Tang, Y.; Goddard, W.A. An Experimental Study of Wetting Behavior and Surfactant EOR in Carbonates with Model Compounds; Society of Petroleum Engineers: Richardson, TX, USA, 2008.


68. Li, Y.V.; Cathles, L.M. Retention of silica nanoparticles on calcium carbonate sands immersed in electrolyte solutions. J. Colloid Interface Sci. 2014, 436, 1–8. [CrossRef] [PubMed]


70. Wenzel, R.N. Resistance of solid surfaces to wetting by water. Ind. Eng. Chem. 1936, 28, 988–994. [CrossRef]


