Interactions of ferro-nanoparticles (hematite and magnetite) with reservoir sandstone: implications for surface adsorption and interfacial tension reduction

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Abstract

There are a few studies on the use of ferro-nanofluids for enhanced oil recovery, despite their magnetic properties; hence, it is needed to study the adsorption of iron oxide (Fe2O3 and Fe3O4) nanoparticles (NPs) on rock surfaces. This is important as the colloidal transport of NPs through the reservoir is subject to particle adsorption on the rock surface. Molecular dynamics simulation was used to determine the interfacial energy (strength) and adsorption of Fe2O3 and Fe3O4 nanofluids infused in reservoir sandstones. Fourier transform infrared spectroscopy and X-ray photon spectroscopy (XPS) were used to monitor interaction of silicate species with Fe2O3 and Fe3O4. The spectral changes show the variation of dominating silicate anions in the solution. Also, the XPS peaks for Si, C and Fe at 190, 285 and 700 eV, respectively, are less distinct in the spectra of sandstone aged in the Fe3O4 nanofluid, suggesting the intense adsorption of the Fe3O4 with the crude oil. The measured IFT for brine/oil, Fe2O3/oil and Fe3O4/oil are 40, 36.17 and 31 mN/m, respectively. Fe3O4 infused with reservoir sandstone exhibits a higher silicate sorption capacity than Fe2O3, due to their larger number of active surface sites and saturation magnetization, which accounts for the effectiveness of Fe3O4 in reducing IFT.

Keywords Magnetite and hematite nanoparticles · Rock surface adsorption · Molecular dynamics simulation · Interfacial tension (IFT)

1 Introduction

The main magnetic molecules in the ferro-nanofluid group are Fe3O4 (magnetite), α-Fe2O3 (hematite, weakly ferromagnetic or antiferromagnetic), γ-Fe2O3 (maghemite, ferrimagnetic), FeO (wineite, antiferromagnetic), ε-Fe2O3 and β-Fe2O3 (Negin et al. 2016; Tsai et al. 2010; Wu et al. 2008). Huh et al. (2015) reported that ferrofluids comprise hydrophobic magnetic, paramagnetic or superparamagnetic iron oxide nanoparticles (NPs). Iron oxide (Fe2O3/Fe3O4) NPs can increase the viscosity of the displacing fluid, resulting in higher sweep efficiency, which is important since most sandstone reservoirs act as neutral or preferentially oil-wet due to adsorption of acidic component of crude oil on their surfaces (Buckley and Liu 1998; Rezaei Gomari et al. 2006). The concept of using ferro-nanofluids was introduced in enhanced oil recovery (EOR)-related processes by Kothari et al. (2010), where they reported that iron oxide NPs are able to reduce viscosity. Kothari et al. (2010) reported that ferro-nanofluids reduce interfacial tension (IFT) in both oil-wet and water-wet reservoirs by causing the collapse of isolated oil bubbles confined in the centre of the pores. Kothari et al. (2010) further hypothesized that the alignment of the reservoir fluid molecules due to the presence of dipole moment reduces flow resistance, possibly leading to higher recovery. Iron oxide NPs have revealed minimal retention in porous media, as shown by a wide-ranging run of transport experiments, which were validated by modelling studies (Yu et al. 2010; Zhang et al. 2015, 2016). However, the research
was limited to the rheological properties of the ferrofluid without investigating the direct application of this fluid in EOR processes.

Haroun et al. (2012) achieved an ultimate recovery of 57% using Fe₂O₃ NPs on carbonate core plugs, while Ogolo et al. (2012) reported an additional recovery up to 24% using Fe₂O₃ NPs dispersed in water. After conducting additional experiments, Ogolo et al. (2012) suggested that iron oxide, when dispersed in brine, performs as a practically good EOR agent in sandstone reservoirs. In addition, Joonaki and Ghanaatian (2014) reported that Fe₂O₃ were only able to recover 17% extra oil, while other NPs (i.e. Al₂O₃ and SiO₂) recovered around 20% additional oil. However, Kamal et al. (2017) suggested that more experimental work is required to understand the underlying mechanism of improvement in interfacial properties using iron oxide NPs. Experimental studies have been carried out in this area largely dealt with determining the optimum NPs concentrations corresponding to minimum IFT. Despite the several studies of the effectiveness of nanofluids in reducing IFT (Hendraningrat et al. 2013; Li et al. 2013; Parvazdavani et al. 2014; Torsater et al. 2012), there is a dearth of research on the potential use of iron oxides for EOR. The mechanisms by which iron oxide (Fe₂O₃ and Fe₃O₄) NPs interact with sandstone to influence reservoir properties are not thoroughly understood (Agista et al. 2018). Given that rock-fluid interactions and IFT play a significant role in oil recovery methods, it is crucial to explore interactions of ferro-nanofluids with reservoir rock to determine the optimum mechanisms for increasing oil recovery. The interfacial reaction between silicates in reservoir system and iron oxide surfaces is of great importance in nanofluid recovery, since IFT and wettability modifications are dependent on adsorption reactions between the ferric species and the silicates in the reservoir sandstone.

NPs are basically used in EOR operations because of their wettability alteration and IFT reduction capabilities (Soleimani et al. 2018). The collision transport of the NPs through the reservoir is subject to particle adsorption on the rock surface, which in turn is controlled by surface charges of the NPs (Dunphy Guzman et al. 2006). In addition, wettability alteration is dependent on adsorption of the nanoparticles on the surfaces of the sandstone grains. Moreover, studies of crude oil adsorption are valuable in directing the oil-displacing agent design and oil exploitation. Hence, there is need to determine possible adsorption as well as the respective adsorption energies of Fe₂O₃ and Fe₃O₄ NPs on the rock surface. Molecular dynamics (MD) simulation was used to determine the interfacial energy (strength) and adsorption configurations of selected ferro-nanofluids (Fe₂O₃ and Fe₃O₄) infused with reservoir sandstones. The density functional theory (DFT) calculation is a multifaceted technique to predict various energetic, structural and electronic properties of diverse systems, thus providing theoretical direction for relevant experimental studies. Spectroscopic techniques such as attenuated total reflection Fourier transform infrared (ATR-FTIR) and X-ray photon spectroscopy (XPS) were used to monitor possible transformations and band shifts in the silicate and C–H species. The adsorption and IFT of Fe₂O₃ and Fe₃O₄ nanofluids were comparatively analysed to determine the better iron species. In general, this study outlines the mechanisms involved in the sorption of Fe₂O₃ and Fe₃O₄ nanofluids onto sandstone surfaces, with respect to EOR.

2 Materials and methods

To evaluate the interactions of ferro-nanofluids with reservoir sandstone, some simulation and experimental studies were carried out. Both approaches are discussed in the underlying subsections.

2.1 Molecular dynamics simulation of adsorption of Fe₂O₃ and Fe₃O₄ NPs on reservoir sandstone surfaces

The adsorption of the selected Fe₂O₃ and Fe₃O₄ NPs in addition to their effects on the shear viscosity of oil was simulated on an atom-based model using the dynamic Forcite tool in the Material Studio software at varying pressures. Molecular dynamics simulation in Material Studio software was applied to simulate the adsorption force between reservoir rock surfaces and molecules of the ferro-nanofluids. Berea sandstone of 24% porosity and bulk composition of quartz, feldspar and kaolinite were used. Specific steps were taken to construct the reservoir sandstone. In step 1, the initial structures of sandstone components (quartz, feldspar and kaolinite) were separately constructed (Fig. 1), and then geometrically optimized to reduce the magnitude of calculated force until they become smaller than defined convergence tolerances (Khaled 2012). An amorphous cell was built. The sandstone components were subsequently imported into the amorphous cell with compositions of 78%, 5%, 5%, 5% and 7% for quartz, anorthite, orthoclase, albite and kaolinite, respectively. In step 2, C₈H₁₈ (clusters of hexane) and brine (H₂O + NaCl) were incorporated into the cell containing the preconditioned mineral surface to closely replicate the reservoir environment (Fig. 1). For step 3, the shared orientation between the oil and the nanoparticle surface was arbitrarily selected, with the oil components placed into a box with same length and width to silica surface. In the course of the simulation, the quartz surfaces were fixed since the vibration of these atoms was minute at room temperature and could be disregarded.

The following scenario involves the adsorption of the ferro-nanofluids on the reservoir sandstone. Prior to the
adsorption configuration, the amorphous cell was transformed into a super cell of \(5 \times 3 \times 1\) with dimensions of \(2.46\) nm \(\times 2.55\) nm \(\times 1.41\) nm and three-dimensional periodic boundary conditions, and a vacuum slab was created afterwards. Similarly, the adsorption comprises specific steps. Firstly, nanofluids of Fe\(_2\)O\(_3\) and Fe\(_3\)O\(_4\) nanoparticles with diameter of 4 Å were built to form the adsorbates. In step 2, the adsorbate was imported to the vacuum slab, and the target atoms were marked (Fig. 2). For step 3, the NPs were placed at the centre of the cubic simulation cell to ensure full dispersion in the reservoir sandstone. A single run from nanoparticle optimization to adsorption took about 72 h. The simulations were performed using the supercell scheme. All NPs and oil chains were fully optimized until the maximum force acting on any atom was lower than 0.001 eV/A.

Due to the presence of the NPs, a universal force field was applied rather than the COMPASS force field. The Materials Studio software comprises a full implementation of the universal force field, including bond order assignment. The adsorption module was then run to create a combined adsorbent–adsorbate structure. The adsorption locator module simulates the reservoir sandstone substrate loaded with the adsorbates of fixed composition. The preferential adsorption sites on the cleaved sandstone surface are targeted by finding the low-energy adsorption sites. The adsorption energies of the nanofluids on the sandstone in the presence of brine and oil were measured using the adsorption configuration locator. The total energy and energy distribution of the adsorption were calculated for the different adsorbates. The energy calculation combines the bonding (stretching, bending, torsion energy and the diagonal and off-diagonal cross-coupling terms) and non-bonding terms. The van der Waals interactions are condensed at \(r_c = 12\) Å by executing a spline function from 11 Å. The Coulomb interactions are calculated via Ewald summation (Allen and Tildesley 1996). The interactions between the NPs and the reservoir sandstone surface are simply intramolecular and van der Waals types. Nonetheless, the intramolecular Coulomb interactions were subjected to Ewald summation in preference to the cut-off method. To properly describe the on-site Coulomb interactions in the localized 3d orbitals of strongly correlated systems, additional Hubbard-type terms were included. All calculations are spin-polarized computations. London dispersion interactions were included in the total bonding energy. Prior to the MD simulations, energy
minimizations were done to relax the local unfavourable structure of the NPs. The surface energy of the oil adsorbed on the sandstone was also determined. The volume of adsorbates was expanded to ensure complete dispersion on the sandstone surface. The stress autocorrelation function (SACF), also referred to as pressure correlation function, was used to estimate the shear viscosity of the reservoir fluid with the incorporation of \( \text{Fe}_2\text{O}_3 \) and \( \text{Fe}_3\text{O}_4 \). Several simulations were performed to fully relax the system, and the trajectories of the final run were selected for analysis.

### 2.2 Stability of interaction: interfacial energy

The binding energy per atom was used as the measure of the stability of the studied systems. It was calculated according to the equation:

\[
E_{b/N} = \frac{1}{N} \left( E_{\text{model tot}} - \sum_{\alpha=1}^{N} E_{\text{atom} \alpha} \right) \tag{1}
\]

where \( N \) is the number of atoms in the system, and \( E_{\text{atom} \alpha} \) is the total atomic energy of the free atom of type \( \alpha \) (H, C, O, Si, Fe). More stable systems are characterized by more negative \( E_{b/N} \).

To further evaluate the strength of interaction between the NPs and oil or water molecules inside pores, the interaction energy was defined. The relative energy of interactions between those components of the system can be described as:

\[
E_{\text{int}} = E_{\text{tot system}} - \sum_{i=1}^{n} E_{\text{tot comp}i} + E_{\text{corr}} \tag{2}
\]

where \( E_{\text{tot system}} \) and \( E_{\text{tot comp}i} \) are the total energies of investigated system and all isolated components in the pore. \( E_{\text{corr}} \) is the...
The interfacial strength is defined as:

\[
\gamma = \frac{E_{\text{model}} - E_{\text{sandstone}} - \sum_{i=1}^{N} E_{\text{comp},i}^\text{cavity}}{A_{\text{cavity}}}
\]

(3)

where \(E_{\text{model}}, E_{\text{sandstone}}, E_{\text{comp},i}^\text{cavity}\) are the total energies of the hybrid model, sandstone and all components in the cavity such as oil and NP, respectively. The interfacial area, \(A_{\text{cavity}}\), is the total surface area of the cavity calculated as solvent-exposed surface area of cavity in sandstone part of the model. High \(\gamma\) value indicates the strong bonding between oil and porous medium, while low \(\gamma\) value suggests increasing separation tendencies in the system. In order to quantify morphology changes of the interface after addition of the chosen nanostructures into the system, the interfacial strength is defined as in Eq. (4).

\[
\gamma = \frac{E_{\text{model}} - E_{\text{sandstone}} - \sum_{i=1}^{N} E_{\text{comp},i}^\text{surface}}{A_{\text{surface}}}
\]

(4)

where \(A_{\text{surface}}\) is the total surface area of the grain calculated as solvent-exposed surface area of quartz grain.

### 2.3 Experimental section

A series of experiments were performed to explore the interactions between iron oxide (Fe\(_2\)O\(_3\)/Fe\(_3\)O\(_4\)) NPs and sandstone in the presence of brine and crude oil. The experiments were conducted using a Berea core sandstone. Brine (NaCl solution) with salinity of 11,000 ppm was prepared to reflect the concentration of infiltrated and connate water (NaCl solution) with salinity of 11,000 ppm was prepared to reflect the concentration of infiltrated and connate water in the offshore sandstone reservoir. The Fe\(_2\)O\(_3\) and Fe\(_3\)O\(_4\) NPs were purchased from Sigma-Aldrich. The NPs (0.5 g each) were dispersed in brine (100 mL) to attain 0.05 wt% of nanofluids. The sandstone core was first aged in the brine for 5 days at 110 °C using the hydrothermal synthesis autoclave reactor, which consists of a double-walled Teflon reaction vessel and has a working pressure of ≤ 3 MPa or 30 bars, to create spontaneous imbibition. Then, a force that correlates the IFT to contact angle, acts on this plate. A vibrating sample magnetometer (VSM) was used to obtain the saturation magnetization of the iron oxide nanoparticles.
3 Results

3.1 Simulation

3.1.1 Adsorption energy

Molecular dynamics simulations are applied to explore the adsorption process of the single- and double-chained magnetite and hematite on SiO$_2$ and other silicate surfaces present in the reservoir sandstone. The total adsorption energy was calculated, which is inclusive of binding, intramolecular and isolated molecular energies. After 10 iterative steps, the adsorption energy for magnetite NPs is about 100 kcal/mol, while that of Fe$_2$O$_3$ NPs is approximately $-2300$ kcal/mol (Fig. 4). The adsorption difference is reflective of surface-structural variations in the adsorbates, adsorbents and interfaces in Fe$_2$O$_3$ and Fe$_3$O$_4$ interactions with the sandstone. The relatively large adsorption energies for both NPs infused with the reservoir sandstone can be attributed to their high molecular weight and reduced long-range diffusion of the inherent hydrodynamic interactions, as well as interaction between the oxygen atoms in Fe$_2$O$_3$ and Fe$_3$O$_4$ compounds and SiO$_2$ surface. The negative energy of Fe$_2$O$_3$ is indicative of exothermic adsorption reaction and implies that the adsorption structures are stable. Here, the adsorption leads to decreased residual forces on the SiO$_2$ surface, which in turn initiates a decline in the surface energy. The negative value essentially indicates physical adsorption. On the other hand, the positive value (+) for Fe$_3$O$_4$ infused with the sandstone indicates an endothermic interaction behaviour, where the intramolecular energy is not dissipated but rather stored in the form of newly created chemical bonds. Thus, the interaction can be considered as chemisorption.

3.1.2 Optimization and interfacial strength

Optimization of the models revealed that Fe$_3$O$_4$ could induce significant changes in the distribution of oil chains around the sand rock surface. In particular, in the presence of Fe$_3$O$_4$ nanoparticles shown in Fig. 5, oil chains are moved or/and bent towards the nanoparticles. The absolute value of sum of the total energies of all isolated components of the systems is bigger than the absolute value of the total energy of the interacting system. In other words, NPs still considerably lowers the interface strength, thus making the removal of oil
(decomposition of the system) energetically preferred. The calculated values of interfacial strength values are given in Table 1. The \( \text{Fe}_3\text{O}_4 \) NPs displayed comparably higher interfacial strength with the sandstone than \( \text{Fe}_2\text{O}_3 \) NPs, as shown in Table 1. However, the interfacial strengths of \( \text{Fe}_3\text{O}_4 \) cluster and \( \text{Fe}_2\text{O}_3 \) NPs are similar, which implies \( \text{Fe}_3\text{O}_4 \) NPs are more effective when stable and well-dispersed in the fluid as compared to when aggregated into clusters.

### 3.1.3 Stress autocorrelation function (SACF)

This pressure function denotes shearing force. The SACF accounts for the stresses imparted on the system due to the diffusion of molecules and intermolecular kinetics, i.e. molecular stresses caused by attraction and repulsion of molecules. However, as it approaches the stability, the SACF starts to converge to a monotonic level, which satisfies that the viscosity analysed is acceptable. The SACF is a protocol under the Forcite module of the molecular dynamics simulations, which was used to calculate shear stresses of \( \text{Fe}_2\text{O}_3 \) and \( \text{Fe}_3\text{O}_4 \) nanofluids at reservoir temperature of 119 °C and pressure range of 500–25,000 psi. The SACF values of the nanofluids across the pressure range (500–25,000 psi) were plotted, as shown in Fig. 6. Interestingly, the SACF values for \( \text{Fe}_2\text{O}_3 \) show consistently higher values of approximately 0.021 across the pressure range; however, the SACF of \( \text{Fe}_3\text{O}_4 \) nanofluid is stable (0.00128) from pressure values of 0–15,000 psi, then increases to 0.025. These SACF values indicate the \( \text{Fe}_2\text{O}_3 \) NPs are more stable within reservoir pressure conditions (4000–5000 psi) and high temperature with suitable viscosity to improve oil mobility and enhance sweep efficiency.

### 3.2 Morphological characterization

#### 3.2.1 Adsorption of \( \text{Fe}_2\text{O}_3 \) and \( \text{Fe}_3\text{O}_4 \) on quartz

The surface morphologies of the reservoir sandstone aged in the ferrofluids was characterized using FESEM. For \( \text{Fe}_2\text{O}_3 \) NPs interaction with sandstone, platy booklets of kaolinite and distinct polygonal quartz crystal faces were observed (Fig. 7a). The characteristic conchoidal and planar fractures of quartz are apparent. The \( \text{Fe}_2\text{O}_3 \) NPs are simply deposited on the quartz surface (i.e. physio-sorption) with no clear solid–solid/liquid–solid phase reaction. In contrast, the \( \text{Fe}_3\text{O}_4 \)–infused reservoir rocks show free-flowing and undulating surfaces with globular-shaped terminations (Fig. 7b), and no tendency to form aggregates. The \( \text{Fe}_3\text{O}_4 \) NPs coat the entire quartz surface, making it indiscernible, indicating occlusion or possible chemisorption of the magnetite on the sandstone.

#### 3.2.2 Elemental mapping

Elemental mapping was conducted on the sandstone surface (specifically Fe ions) to corroborate the FESEM micrographs. The Fe elements are sparsely distributed across the surface of \( \text{Fe}_2\text{O}_3 \) imbued with sandstone (Fig. 7c). In contrary, the mapped FESEM image of \( \text{Fe}_3\text{O}_4 \) NPs infused with reservoir sandstone show uniform distribution of Fe element over the sandstone surface (Fig. 7d). This disparity in Fe distribution further suggests the higher adsorption capacity of the \( \text{Fe}_3\text{O}_4 \) nanofluid on the reservoir sandstone compared to \( \text{Fe}_2\text{O}_3 \). This higher dispersibility of \( \text{Fe}_3\text{O}_4 \) supports its ability to infuse with oil and disrupt the wettability of the oil molecules on the rock surface. The images and EDX analysis of the sandstone pore (Fig. 7d) indicate the presence of Fe infused with the oil and matrix material, which infers that the \( \text{Fe}_3\text{O}_4 \) effectively infiltrates the rock and adsorbs on the pore filling matrix material.

#### 3.2.3 EDX composition analysis

The elemental concentrations of the ferro-nanofluids infused with the reservoir sandstone were analysed with EDX (Table 2). Spot analysis was performed for the sandstone grain and matrix (pore infill) material. The absorbed

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**Table 1** Interfacial strengths (\( \gamma \)) of systems containing quartz grains, three oil chains and different types of nanostructures

| Grain  | Nanostructure         | Interfacial strength \( \gamma \), J/m² |
|--------|-----------------------|----------------------------------------|
| Quartz | \( \text{Fe}_2\text{O}_3 \) (\( r_{in} = 5 \text{ Å} \)) | 0.06797                               |
| Quartz | \( \text{Fe}_3\text{O}_4 \) (\( r_{in} = 5 \text{ Å} \)) | 0.12538                               |
| Quartz | \( \text{Fe}_3\text{O}_4 \) (cluster) | -0.005                                |

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**Fig. 6** Plot of SACF values for \( \text{Fe}_2\text{O}_3 \) and \( \text{Fe}_3\text{O}_4 \) nanofluids at different pressures
Fe ions were found to vary between the infused Fe₂O₃ and Fe₃O₄. The Fe₂O₃ infused with sandstone showed relatively lower adsorbed Fe ions (≈0.5%) compared to Fe₃O₄ infused with sandstone (≈2%). The amount of adsorbed carbon also varied from 50% for Fe₂O₃ infused sample to 80% for Fe₃O₄ infused sample. This implies that the concentration

**Fig. 7** FESEM micrographs of a Fe₂O₃ infused with reservoir sandstone, with inset arrow indicating quartz fracture and b Fe₃O₄ infused with reservoir sandstone, with the inset arrow showing quartz terminations are coated. Surface Fe elemental mapping of c Fe₂O₃, and d Fe₃O₄ infused with reservoir sandstone

| Sample                  | S (wt%) | O (wt%) | Fe (at%) | C (wt%) | Al (wt%) | Na (wt%) | Fe (wt%) | Cl (wt%) |
|-------------------------|---------|---------|----------|---------|----------|----------|----------|----------|
| Fe₂O₃ infused with sandstone | 17.14   | 7.93    | 37.21    | 42.82   | 54.30    | 0.59     | 0.22     | 1.64     |
|                         | 39.57   | 30.07   | 43.29    | 57.73   | –        | 1.97     | 5.63     | 1.34     |
|                         | 11.31   | 9.43    | 31.99    | 45.57   | –        | 10.38    | 22.28    | 1.52     |
| Fe₃O₄ infused with sandstone | 13.83   | 7.34    | 6.59     | 60.59   | 76.11    | 3.14     | 8.49     | 7.55     |
|                         | 15.77   | 9.01    | 13.48    | 62.77   | 75.66    | –        | 0.95     | 6.96     |
|                         | 3.43    | 1.67    | 15.55    | 71.69   | 81.64    | 2.43     | 0.38     | 6.53     |

Table 2: Elemental distributions (weight percent (wt%) and atomic percent (at%)) of Fe₂O₃ and Fe₃O₄ infused with sandstone, showing higher adsorbed Fe ions and carbon in the case of Fe₃O₄.
of available C atoms from the oil retained in the reservoir system increases with adsorbed Fe.

3.3 FTIR analysis

The FTIR spectra of the various sandstones aged under different conditions (brine, brine + crude, brine + crude + nano-fluids) are shown in Figs. 8, 9 and 10. The functional groups (band assignments) detected by FTIR spectroscopy of the sandstone samples are given in Table 3. The functional groups detected in the FTIR spectra of the various samples are presented and their variations are determined in the underlying subsections.

3.3.1 Silicate and H–O–H bonds

The FTIR spectra of the sandstone samples show the characteristic peaks for siloxane (Si–O–Si), Si–O quartz, Si–O stretching and bending vibrations (Fig. 8). After ageing with crude oil, brine, and Fe$_2$O$_3$ and Fe$_3$O$_4$ nanofluid solutions, the characteristic peaks of Si–O symmetrical stretching, Si–O symmetrical bending and Si–O asymmetrical stretching vibrations within the ranges of 694, 777–797 and 1083–1099 cm$^{-1}$, respectively, remain for all samples (Fig. 8). However, the absorbance peaks for Si–O–Si bending at 644 cm$^{-1}$ and H–O–H bending of water 1640 cm$^{-1}$ in the spectra of sandstone aged in brine, but absent in samples infused with oil, indicate the masking of the siloxane bond and displacement of water by crude oil. The distinct absorption bands at 1030 cm$^{-1}$ and 1006 cm$^{-1}$ are ascribed to framework Si–O–Si stretching vibrations. The absorption bands at 935 cm$^{-1}$ and 910 cm$^{-1}$ represent the Al–OH bending vibrations. The band at 935 cm$^{-1}$ is also attributed to the non-bonding of inner surface or surface (nanostructure) OH. The reduction in the cusp of the characteristic peak of H–O–H bonding in the spectrum for sandstone immersed in brine can be attributed to the ability of Fe$_2$O$_3$ and Fe$_3$O$_4$ fluids to accelerate oil removal in water–oil emulsion (Ko et al. 2014). The observed shift in the siloxane (Si–O–Si) and Si–O quartz bonds indicate adsorption of Fe–O from magnetite to SiO$_2$.

Higher-resolution FTIR spectra were obtained for low-wave number region (1300–650 cm$^{-1}$) to clearly show the variations in the functional groups (Fig. 9). The characteristic absorption bands for Fe(hydr)oxides were identifiable within the spectral range of 170 cm$^{-1}$ and 570 cm$^{-1}$, which were beyond the detection limit of instrument used in this study. Absorption bond was identified at 750 cm$^{-1}$ in FTIR spectra of Fe$_3$O$_4$ infused in the sandstone, which is assigned to out of plane FeOOH vibrations. Nonetheless, the Fe–O bond indicates the presence of maghemite, a defective form of magnetite. The presence of FeOOH and Fe–O

![Fig. 8 Stacked FTIR spectra of sandstone, sandstone aged in crude oil, sandstone aged in crude oil and Fe$_2$O$_3$ nanofluid, and sandstone aged in crude oil and Fe$_3$O$_4$ nanofluid](image-url)
Fig. 9 Closer view of the low-wave FTIR region (600–1300 cm$^{-1}$) for sandstone, sandstone aged in crude oil, sandstone aged in crude oil and Fe$_2$O$_3$ nanofluid, and sandstone aged in crude oil and Fe$_3$O$_4$ nanofluid

Fig. 10 Overlapping FTIR absorbance spectra of sandstone, sandstone aged in crude oil, sandstone aged in crude oil and Fe$_2$O$_3$ nanofluid, and sandstone aged in crude oil and Fe$_3$O$_4$ nanofluid
bands affirms the existence of both ferrous (Fe²⁺) and ferric (Fe³⁺) ions, which implies the stability of the Fe₃O₄ within an environment where oxidative and reductive states of Fe occur and are sustained.

### 3.3.2 C–H bonds

Two intense C–H stretching vibrations were also identified at around 2920 cm⁻¹ and 2850 cm⁻¹, for sandstone samples saturated with crude oil, brine, and nanofluid solution. The C–H stretching bands in 2850–2958 cm⁻¹ are indicative of polyatomic Cₙ–H–O entities with C bonded to two or three H. The highest intensity (vCH) band between 2920 and 2931 cm⁻¹ is assigned to symmetrical stretching of C–H mode of –CH₃ group. A spectral peak is also discernible at 1460 cm⁻¹, which is assigned to C–H deformation of the saturate due to vibration of the carboxylic acid group present in the crude. C–H symmetric deformation of the saturate (1376 cm⁻¹) was also detected, although with larger intensity in the sample aged in the Fe₃O₄ nanofluid. This relatively higher intensity is suggestive of the displacement of oil by the Fe₃O₄ nanofluid, thus enabling improved detection of the C–H peak. In addition, the presence of a peak denoting substituted aromatics C–H bonds of benzene ring at 715 cm⁻¹ (Fig. 8), which is conspicuously absent in the spectra of other samples. This band is assigned to long-chain methyl rock. Stacked FTIR spectra show variations in peaks and band intensities. The FTIR spectrum for sandstone samples treated with crude oil and Fe₃O₄ shows the highest peak intensities for characteristic C–H bond stretching (symmetric and asymmetric) vibration. Overlapping FTIR spectra show characteristic C–H bond stretching (symmetric and asymmetric) vibration for sandstone samples treated with crude oil, and Fe₃O₄ nanofluid have the highest peak compared to the other samples. The saturate fraction has dominant aliphatic hydrocarbons (alkanes) as indicated by the complete presence of absorbance near 2920, 2850 and 1450 cm⁻¹ and the absence of absorption near 1600 cm⁻¹, which is the characteristic band for aromaticity. Although the presence of the absorption band at 715 cm⁻¹ in the spectra of Fe₃O₄ infused

### Table 3  Functional groups (band assignments) detected by FTIR spectroscopy of the sandstone samples

| Wave number, cm⁻¹ | Type of vibration                                                                 |
|-------------------|-----------------------------------------------------------------------------------|
| 3640              | Al–O–H stretching, OH stretching, crystalline hydroxyl                           |
| 3418              | Stretching modes of surface H₂O molecules or envelope of hydrogen-bonded (absorbed) surface OH groups |
| 2916              | Asymmetric and symmetric stretching vibrations of C–H bonds of aliphatic CH₃ group |
| 1644              | Asymmetric and symmetric stretching vibrations of C–H bonds of aliphatic CH₃ group |
| 1388              | C–H deformation (bending vibration of C–H of CH₂ and CH₃ aliphatic groups)         |
| 1028              | C–H symmetric deformation                                                          |
| 1001              | Si–O asymmetrical stretching vibration; clay minerals                               |
| 1084              | Si–O stretching vibration                                                           |
| 935               | Si–O quinze                                                                           |
| 912               | Non-bonded inner surface or surface (nanostructure) OH; Si–O stretching vibration    |
| 795               | Al–OH deformation vibration                                                        |
| 776               | Si–O symmetrical stretching vibration                                               |
| 735               | Si–O quartz hydroxyl group (Al–OH) perpendicular to the surface (Translational –OH) |
| 692               | Substituted aromatics C–H bonds of benzene ring                                    |
| 644               | Si–O symmetrical bending vibrations                                                |
with the sandstone sample is proof of the aromaticity, the C–H stretch band at 3000–2850 cm$^{-1}$ is the most intense because the change in dipole moment with respect to distance for the C–H stretching is greater compared to others. The weak bond at 3083 cm$^{-1}$ is assigned to =C–H stretch, which indicates the presence of alkenes and aromatics.

The Fe$_3$O$_4$ infused sandstone sample shows the highest absorbance for the Si–O quartz, Si–O–Si and C–H bonds (Fig. 10). The higher intensity for Fe$_3$O$_4$-infused sandstone sample is attributed to greater change in dipole moment with respect to distance. The peak shifts and higher adsorbance intensities in the case of Fe$_3$O$_4$-infused sandstone sample imply disturbance of the adsorption layers of oil molecules existing on the solid surface by FeOOH and Fe–O compounds, which contributes to the detached oil molecules. The cusp observed for the pure sandstone, which is absent in the other spectra, denotes stretching modes of surface H$_2$O molecules. This absence suggests the possible expulsion of connate and adsorbed water molecules with the introduction of the NPs.

### 3.4 X-ray photon spectroscopy

The powder samples were also studied with XPS to comprehensively analyse the implications of their oxidative phase, which determines the properties of iron oxide NPs. The XPS method subjects the samples to EM irradiation from a soft X-ray source under ultrahigh vacuum, which results in the absorption of the EM by the electrons contained in the samples (with binding energy less than the vacuum level). The absorbed EM radiation is subsequently dispelled from the solid with a kinetic energy that is then analysed. The scattered electrons have an energy that is typical of the element from which they are emitted, and the intensity of the characteristic peaks observed in the energy spectra is related to the concentration of the elements present in the surface through relative sensitivity factors proper of each instrument. The relative intensities of the oxides and elemental peaks change with the oxide thickness.

XPS spectra were obtained for different samples (Fig. 11). The spectra reveal the existence of carbon, sodium (Na 1$s$, 1071–1072.5 eV; derived from the brine used to disperse the nanoparticles), oxygen (1$s$, 532.2 eV), two forms of carbon: carbon singly bonded to an oxygen atom (1$s$, 286.5 eV, and carbon doubly bonded to a hydrogen atom as in aliphatic chain (1$s$, 284.6 eV), two forms of silicon: silicon singly bonded to an oxygen atom (2$p$, 103.5 eV) and silicon bonded to a C–H chain (2$p$, 102.5 eV), aluminium (2$p$, 74.5 eV) and Fe (accessory minerals in the Berea sandstone). Some samples also exhibit chlorine (2$p$, 198.5 eV). The reference line for all the XPS spectra was organic carbon (1$s$, 284.6 eV). Adventitious carbon content is negligible. The strong intensity peak of C suggests organic/biogenic source or catalytic interaction of C- and H-containing compounds and the quartz surface, rather than from mere exposure to air. The C centred at 285 eV denotes organic species with C–H, C–C and C–O bonds (Hochella 1988; Hochella et al. 1990). The existence of the oxygenated carbon may possibly augment the hydrophilic character of the ferrofluids. However, the characteristic XPS peaks for Si, C and Fe are absent in the spectral of sandstone samples aged in magnetite, suggesting the intense adsorption of the magnetite with the crude oil. Underlying subsections provide detailed explanation of the XPS survey spectra for silicon and carbon. The XPS parameters are given in Table 4.

#### 3.4.1 XPS silicon spectra

The chemical state of Si in these XPS spectra is SiO$_2$ (quartz), as indicated by the binding energies at $\approx$ 104 eV. The spectral variation of the different samples with regard to Si is shown in Fig. 12. Fe$_3$O$_4$ pure sandstone (Fig. 12a), the identified peaks include Si 2$p$ (oxide) and Si 2$p$ (element) at 104 eV and 103.5 eV, respectively. With the addition of crude oil and Fe$_3$O$_4$, the characteristic peak for Si 2$p$ (oxide) peak becomes symmetric at binding energy of $\approx$ 104 eV (Fig. 12b, c). For Fe$_3$O$_4$, the characteristic peak for Si 2$p$ (oxide) diminishes. This distorted binding energy spectrum for Fe$_3$O$_4$-infused sample (Fig. 12d) suggests high adsorption and interfacial energy between Si and Fe$_3$O$_4$, which led to the poor resolution of Si 2$p$-orbit components.

#### 3.4.2 XPS carbon spectra

For the bare sandstone sample (Fig. 13a), the carbon identified at binding energy of $\approx$ 285.7 eV denotes adventitious carbon (C–O–C), which is likely derived from exposure of the sample to the atmosphere. Also observed is a C 1$s$ component at the left, indicating the presence of ester (O–C=O), which similarly points to adventitious carbon. For the sample aged with crude oil and brine, the carbon peak is identified at energy position of 286 eV with high counts/s of greater than 17,000 (Fig. 13b, c). Based on its symmetry, this C 1$s$ peak is denoted as phenyl. Extended delocalized electrons in the sample (e.g. aromatic rings) can result in a satellite structure, several eV to higher binding energy of the main peak, as in the case of Fe$_3$O$_4$-infused oil (Fig. 13d).

#### 3.5 Interfacial tension (IFT)

The interfacial tensions between crude oil and nanometric colloidal Fe$_2$O$_3$ and Fe$_3$O$_4$ dispersed in brine were investigated. The measured IFT for brine/oil, Fe$_2$O$_3$/oil and Fe$_3$O$_4$/oil are 40, 36.17, and 35 mN/m, respectively. Figure 14 shows the distribution of IFT values with time for different nanofluids and brine as control sample. The IFT
properties are dependent on the particle attachment at the interface. The variation in IFT is as a result of the hydrophilic or lipophilic character of the particles, which is the basis of their affinity for the fluid interface. For the Fe$_2$O$_3$ nanofluid infused with crude oil, the utilized particles are strongly hydrophilic, which are then completely wetted and do not influence the properties of the aqueous dispersion interface. Instead, the surfaces of Fe$_3$O$_4$ nanoparticles have a higher hydrophobicity and adsorption, providing a driving force for their attachment at the biphasic liquid/fluid interface (Felicia et al. 2016). Thus, Fe$_3$O$_4$ reduced IFT by as much as approximately 14.3%, whereas Fe$_2$O$_3$ showed a slightly lower reduction of 3.3% on the average. The reduced IFT can be attributed to the electrostatic repulsive interactions between the Fe$_3$O$_4$ NPs and the oil surface, which promotes the diffusion of the nanoparticle towards the interface (Ma et al. 2008). The nanoparticle adsorption strongly influences the hydrophobicity/

Fig. 11 XPS survey spectra of a sandstone, b sandstone aged in crude oil, c sandstone aged in crude oil and Fe$_2$O$_3$ nanofluid, d sandstone aged in crude oil and Fe$_3$O$_4$ nanofluid
lipophilicity of the particle, i.e. the particle affinity for the fluid interfacial environment (Levine et al. 1989). The Brownian transport of particles from the bulk to the interface, argued on the basis of the relaxation of dynamic interfacial tension after large expansions of the interfacial area, is most likely ineffective here. In fact, for energetic reasons, the attachment of the particles at the fluid interface is essentially an irreversible process. Moreover, the strongly magnetoresponsive character of $\text{Fe}_3\text{O}_4$ accounts for its high adsorption energy (Socoliuc and Vékás 2014).

### 3.6 Magnetization measurements

The magnetic properties of the $\text{Fe}_2\text{O}_3$ and $\text{Fe}_3\text{O}_4$ NPs were also studied using VSM. The hysteresis loops of $\text{Fe}_2\text{O}_3$ and $\text{Fe}_3\text{O}_4$ NPs at room temperature are shown in Fig. 15. The magnetization study conducted as a function of magnetic field shows hysteresis behaviour with coercivity ($G$) of 123.61 and 96.24 for $\text{Fe}_2\text{O}_3$ and $\text{Fe}_3\text{O}_4$, respectively. The remanent magnetization (retentivity) values are 3.56 and 4.013 emu/g, and the saturation magnetization values are 32.25 and 40.97 emu/g for $\text{Fe}_2\text{O}_3$ and $\text{Fe}_3\text{O}_4$, respectively (Table 5). The higher saturation magnetization and retentivity for $\text{Fe}_3\text{O}_4$ can be ascribed to its greater interparticle magnetic interaction. However, the value obtained for magnetite in this study is much smaller than 91.5 emu/g reported for nanocrystalline magnetite (Ozdemir et al. 2002), and 98 emu/g recorded for bulk magnetite (Cullity 1972). That significant incongruity can be attributed to a possible nonparallel coupling of the magnetization in the surface regions of grains or the presence of impurities that might have appeared during synthesis (Starowicz et al. 2011). The nanoparticle size could also account for this relatively low magnetization.

### 4 Discussion

Given the dearth of knowledge on the application of ferrofluids in EOR, this study investigated the interaction of $\text{Fe}_2\text{O}_3$ and $\text{Fe}_3\text{O}_4$ NPs with sandstone. The kinetics of sorption seems to be faster for $\text{Fe}_3\text{O}_4$ than for $\text{Fe}_2\text{O}_3$. Based on its high adsorption to the reservoir sandstone and ability to infill pore spaces, the effect of $\text{Fe}_3\text{O}_4$ on hydrocarbon recovery is twofold. The $\text{Fe}_3\text{O}_4$ can either increase the viscosity of the displacing fluid, resulting in a higher sweep efficiency or infuse (soaked) with the oil, which can then be attracted and recovered through the application of inward magnetic forces towards the borehole. The SACF value of $\text{Fe}_3\text{O}_4$ corroborates this ability to accelerate oil removal in a water-in-oil emulsion. Thickening of the $\text{Fe}_3\text{O}_4$ infused with crude oil occurs due to ability of the NPs to form a network structure via hydrogen bonding (Zeyghami et al. 2014), which will directly affect the fluid shear stress (Cheraghian and Hendraningrat 2016). On the other hand, the negative charges on the FeOOH surface of $\text{Fe}_2\text{O}_3$ are higher, which accounts for the less favourable adsorption. The results show that silica and clays exhibit a high affinity to surfaces of $\text{Fe}_3\text{O}_4$, as confirmed by the significant increase in absorbance spectral intensity. It is therefore reasonable to suggest that the increase in intensity is due to the sorption of low polymeric silicate species and possibly the enhanced polymerization at the $\text{Fe}_3\text{O}_4$ surface. Thus, the strong infrared absorbance in the wavenumber region 1050–1200 cm$^{-1}$ is related to the polymeric species, while the appearance of the band at about 1120 cm$^{-1}$ could be an infrared spectral evidence for the

### Table 4 XPS parameters for $\text{Fe}_2\text{O}_3$ and $\text{Fe}_3\text{O}_4$ infused with reservoir sandstone

| Sample                   | Si 2p bending energy, eV | Si 2p counts/s | C 1s bending energy, eV | C 1s counts/s | Fe 2p$_{3/2}$ bending energy, eV | Fe 2p$_{3/2}$ counts/s | Fe 2p$_{1/2}$ bending energy, eV | Fe 2p$_{1/2}$ counts/s |
|--------------------------|--------------------------|----------------|-------------------------|--------------|-------------------------------|------------------------|-------------------------------|-------------------------|
| Sandstone                | 104                      | 1100           | 285.7                   | 5372         | 713                           | 30,218.5               | 724                           | 29,589                  |
| Sandstone + crude oil    | 105                      | 400            | 286                     | 16,983       | 712                           | 17,150                 | 724                           | 17,106                  |
| Sandstone + crude oil + $\text{Fe}_2\text{O}_3$ | 103                      | 1400           | 285.5                   | 17,096       | 710                           | 26,446                 | 723                           | 26,350                  |
| Sandstone + crude oil + $\text{Fe}_3\text{O}_4$ | 103.5                    | 79             | 285.6                   | 16,380       | 709                           | 11,464                 | 725                           | 11,262                  |

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formation of a 3D framework silica structure on the Fe$_3$O$_4$ surface.

The simulated and measured parameters for Fe$_2$O$_3$ and Fe$_3$O$_4$ ferrofluids are given in Table 6. Fe$_2$O$_3$ and Fe$_3$O$_4$ ferrofluids were shown experimentally to reduce IFT, with Fe$_3$O$_4$ ferrofluid presenting a higher decrease. By decreasing the IFT and altering the rock wettability using ferrofluids, the capillary pressure will decline, resulting in improved oil recovery (Roustaei et al. 2012). In the case of Fe$_2$O$_3$, the self-organization of dipolar Fe$_2$O$_3$ cubes into dipolar chain morphology accounts for its effect on IFT. Fe$_3$O$_4$ NPs apparently improved the rheology of the dispersing brine solution as it adsorbed to the fluid. The Brownian motion of the Fe$_3$O$_4$ NPs in the bulk and the internal restructuring of the

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**Fig. 12** High-resolution XPS spectra of silicon (Si 2p) for a sandstone, b sandstone aged in crude oil, c sandstone aged in crude oil and Fe$_2$O$_3$ nanofluid, and d sandstone aged in crude oil and Fe$_3$O$_4$ nanofluid
mixed particle (brine-Fe$_3$O$_4$-oil) layer at the interface and the Fe$_3$O$_4$ rearrangement between the attached nanoparticles and the fluid interface (nanofluid solution/hexane interface) account for the IFT reduction (Ravera et al. 2006).

The high adsorption property of Fe$_3$O$_4$ also supports its potential use as a removal agent of microscopic oil droplets and remnant polymer from produced water, as well as divalent cations (Ca$^{2+}$) from hard brine. The catalytic properties of Fe$_3$O$_4$ NPs can potentially enable in situ cracking of oil chains, significantly improving the efficiency of EOR. Thus, the magnetite normally exerts effects on oil degradation by modifying the concentrations of the free radicals. This study is consistent with the previous studies. Shekhawat et al. (2016) proposed the application of Fe$_3$O$_4$ nanofluids for magnetic heavy oil recovery. The differential impact of the ferro-nanofluids is compelled by competing anisotropic interactions caused by the variable particles’ shapes (spherical (Fe$_3$O$_4$) and

![Fig. 13 High-resolution XPS spectra of carbon (C 1s) for a sandstone, b sandstone aged in crude oil, c sandstone aged in crude oil and Fe$_2$O$_3$ nanofluid, and d sandstone aged in crude oil and Fe$_3$O$_4$ nanofluid](image-url)
cubic \([\text{Fe}_2\text{O}_3]\) and their dipole moments (Rossi et al. 2018). As indicated by their saturation magnetization and retentivity (remanent magnetization), the \(\text{Fe}_3\text{O}_4\) displays higher interparticle magnetic interactions, which enables interaction between the oil and \(\text{Fe}_3\text{O}_4\).

### 5 Conclusion

In situ ATR-FTIR and XPS spectroscopies were used to monitor the sorption of ferro-nanoparticles (\(\text{Fe}_2\text{O}_3\) and \(\text{Fe}_3\text{O}_4\)) on silicates. The observed spectral changes indicate \(\text{Fe}_3\text{O}_4\) infused with the reservoir sandstone has a higher silicate adsorption capacity than \(\text{Fe}_2\text{O}_3\), possibly because of their larger number of active surface sites and the existence of both ferrous (\(\text{Fe}^{2+}\)) and ferric (\(\text{Fe}^{3+}\)) ions, which implies the stability of the magnetite within an environment where oxidative and reductive states of Fe are present. The \(\text{Fe}_3\text{O}_4\) nanofluid exhibits a higher IFT reduction, which can be attributed to its strongly magnetoresponsive characteristics, higher hydrophobicity and adsorption, providing a driving force for their attachment at the biphasic liquid/fluid interface.
### Table 6 Values of simulated and measured parameters for Fe$_3$O$_4$ and Fe$_2$O$_3$ infused with reservoir sandstone

| Simulated and measured parameters | Fe$_3$O$_4$ infused with reservoir sandstone | Fe$_2$O$_3$ infused with reservoir sandstone |
|-----------------------------------|--------------------------------------------|--------------------------------------------|
| Adsorption, kcal/mol              | $-2300$                                     | $100$                                      |
| SACF                              | $0.021$                                     | $0.00128$                                  |
| Interfacial strength $\gamma$, J/m$^2$ | $0.06797$                                | $0.12538$                                  |
| Interfacial tension, mN/m         | $36.17$                                    | $35$                                       |
| Adsorbed Fe ions, %               | $0.5$                                      | $2$                                        |

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