Effect of SiO₂ on the foamability, thermal stability and interfacial tension of a novel nano-fluid hybrid surfactant

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ABSTRACT

In foam assisted water alternating gas (FAWAG) method, thermal stability, foamability and interfacial effect of foaming agents (surfactants) are important properties governing the success of oil recovery. The effect of high reservoir temperature is detrimental to these properties, which becomes a great challenge for the applications of surfactants in enhanced oil recovery (EOR). Sικα nano-fluid (SiO₂), which is a recently employed technology, has been utilized to improve the rheology, stability and interfacial properties of surfactants. This study is aimed at investigating the effect of SiO₂ in improving the thermal stability, relative foamability and interfacial (IFT) effect of industrial based surfactant (IBS). Design Expert Software (DOE) using central composite design (CCD) at five levels (-1.68 to +1.68) was employed in the experimental design. The chemical interaction between the SiO₂ and IBS in the novel nano-fluid hybrid surfactant (SiO₂-IBS) had been successfully established using different spectroscopic instruments (FTIR, XRD, FESEM etc.). Furthermore, under the optimum conditions established, SiO₂ had significant effects on the relative foamability and thermal stability of the hybrid material (SiO₂-IBS) at 25-110 °C, and their synergistic effect had been quantified in the multivariate models (cubic). However, the IFT result indicated that presence of the SiO₂ in the hybrid had reduced the IFT drastically from 120.3 ± 9.8 mN/m to 10.6 ± 6.8 mN/m. Consequently, the novel SiO₂-IBS nano hybrid surfactant could be a suitable flooding agent in the high temperature reservoirs for application in FAWAG method.

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1. Introduction

Enhanced Oil Recovery (EOR), an advanced technology employed in oil industry, has the potential to extract residual oil after primary and secondary technologies (Ahmadi et al., 2015; Hamza et al., 2017). The advantage of EOR over the conventional technologies (primary and secondary) involves the injection of foreign fluids (surfactants, polymers, gases etc.) or energy capable of interfering with oil/water/rock systems to mobilize the oil to the surface (Alagorni et al., 2015). Approximately, 40 to 60 % of oil production rate has been attained with the utilization of the EOR technologies (Muggeridge et al., 2014). Water Alternating Gas (WAG) technology, which requires simultaneous injection of gas and water into the reservoir has been the most frequently employed EOR technology (Riazi and Golkari, 2016). However, early gas breakthrough, gravity override and viscous fingering constituted the main challenge in the WAG execution, and result to poor sweep efficiency over time (Qureshi et al., 2016). Efforts to overcome these problems had led to the development of foams to assist in WAG, and this technology is referred to as Form Assisted Water Alternating Gas (FAWAG). In FAWAG, the gas is injected into surfactant solution, which is then trapped to form foam bubbles, these help to control the gas mobility and improve oil recovery (Shabib-asl et al., 2014, Tunio and Chandio, 2012). Despite high oil recovery achieved in FAWAG, obtaining stable foams in high temperature oil reservoirs are difficult and consequently affect their overall performance (Kapetas et al., 2016). However, silica nanofluid (SiO₂), which is a recently employed technology, has been found to show a remarkable...
improve the rheology, stability and interfacial properties of surfactants (Esmaeeli Azadgoleh et al., 2014; Sharma et al., 2016). The effects of SiO2 on stabilizing foams in different types of surfactants have been reported (Xue et al., 2016; Sun et al., 2015). In this study, SiO2 and industrial based surfactant (IBS) were used to formulate a new nanofluid hybrid surfactant (SiO2-IBS), and the effect of the SiO2 on the foamability, thermal stability and interfacial tension of SiO2-IBS had been investigated. The Design Expert Software (DOE) has been used to develop experimental combinations with respect to the concentrations of SiO2 and IBS, as well as aging. The central composite design (CCD) and response surface methodology (RSM) were applied for optimization and modeling.

2. Materials and methods

The SiO2 was purchased as a bare nano powder (20-30 nm, 99.5 wt %, amorphous) from US Research nanomaterials, Inc. USA. It has the surface area of 180-600 m2/g, bulk density and true density of < 0.10 and 2.4 g/cm3, respectively. While the surfactant (IBS), was supplied from a Malaysian local oil company. The compositions of synthetic brine (3.5 %) used is presented in Table 1.

Table 1: Synthetic brine composition

| S/N | Salt name          | Chemical formula |
|-----|--------------------|------------------|
| 1   | Sodium chloride    | NaCl             |
| 2   | Potassium chloride | KCl              |
| 3   | Magnesium chloride | MgCl2·6H2O       |
| 4   | Calcium chloride   | CaCl2·2H2O       |
| 5   | Sodium bicarbonate | NaHCO3           |
| 6   | Sodium sulfate     | Na2SO4           |
| 7   | Strontium chloride | SrCl2·6H2O       |

2.1. Development of combinations

The DOE software was employed to statistically design the formulation combinations using a CCD. Table 2 shows the range values of SiO2, IBS and aging chosen at five levels from -1.68 to +1.68. The choice of this range was because of the conflicting levels of optimum conditions for SiO2, IBS and aging from the literature.

Table 2: Processed factors of independent variables at five levels intervals

| Independent variables | Process level of variables at five levels intervals (α to +α) |
|-----------------------|---------------------------------------------------------------|
| SiO2 (%)              | -1.68 -1 0 +1 +1.68                                         |
| IBS (%)               |                                                             |
| Aging (days)          |                                                             |

The actual processed factors of all the input variables were keyed into the software and Table 3 was generated. The table described 17 different formulation combinations with 6 central points (formulations 15-20) as replicates to minimize error according to CCD. These had let to the development of 20 experimental combinations formulated.

Table 3: Experimental combinations of input variables (in terms of processed factors)

| S/N | Standard Experiments | Factor A IBS | Factor B SiO2 | Factor C Aging |
|-----|----------------------|--------------|---------------|----------------|
| 1   | S1                   | 0.00         | -1.68         | 0.00           |
| 2   | S4                   | 1.00         | 1.00          | -1.00          |
| 3   | S7                   | -1.00        | 1.00          | 1.00           |
| 4   | S2                   | 1.00         | -1.00         | -1.00          |
| 5   | S8                   | 1.00         | 1.00          | 1.00           |
| 6   | S16                  | 0.00         | 0.00          | 0.00           |
| 7   | S15                  | 0.00         | 0.00          | 0.00           |
| 8   | S9                   | -1.68        | 0.00          | 0.00           |
| 9   | S5                   | -1.00        | -1.00         | 1.00           |
| 10  | S13                  | 0.00         | 0.00          | -1.68          |
| 11  | S6                   | 1.00         | -1.00         | 1.00           |
| 12  | S17                  | 0.00         | 0.00          | 0.00           |
| 13  | S1                   | -1.00        | -1.00         | -1.00          |
| 14  | S18                  | 0.00         | 0.00          | 0.00           |
| 15  | S3                   | -1.00        | 1.00          | -1.00          |
| 16  | S20                  | 0.00         | 0.00          | 0.00           |
| 17  | S19                  | 0.00         | 0.00          | 0.00           |
| 18  | S14                  | 0.00         | 0.00          | 1.68           |
| 19  | S12                  | 0.00         | 1.68          | 0.00           |
| 20  | S10                  | 1.68         | 0.00          | 0.00           |

2.2. Chemical formulations

In EOR process, the low concentrations of injecting chemicals are important to achieve high recovery factor at low costs to maximize profit (Han et al., 2013). In this study, the concentrations of the SiO2 and IBS in brine are in the range of -1.68 to +1.68 according to Table 2. These solutions were then combined to make up 60 ml of formulations, as described in the Table 3. All the chemical formulations prepared were stirred at 25 ºC for 12 h using a magnetic stirrer (IKA C-MAG HS 7 S002) set at 600 rpm. Thereafter, the formulations were sonicated at 25 Hz in an ultrasonic bath (Ultrasonics Corporation, USA) for 1 h to minimize particles agglomeration. They were then kept at 25 ºC for aging.

2.3. Compatibility and stability study

Approximately, 60 ml of SiO2-IBS formulations as shown in Fig. 1 were visually observed for 24 h at 25 and 100 ºC, respectively, to find out the compatibility between the SiO2 and IBS. The compatibility codes of A: homogenous solutions and B: precipitates were used to assess the compatibility studies. Furthermore, Fourier Transform Infrared Spectroscopy (FTIR) was conducted to establish the chemical interactions between the hybrid components as evidence of compatibility. For stability studies, four different experiments which include; pH control, viscosity enhancement, ultrasonication and particle size were conducted as described in (Ghadimi et al., 2011). After the normal pH of the solutions were recorded, the pH were then adjusted to 4, 6, 8, 10 and 12 using 1 N HCl and KOH, respectively, and visually observed at an interval of...
time. Other fresh samples were prepared for stability test by viscosity enhancer method. Here a water soluble polymer, polyacrylamide (PAM) was used to increase the viscosity of base fluids in order to decrease the particle’s speed to improve the stability. The concentrations of PAM used were 0.05, 0.1 and 0.15 % in the hybrid, while the controls (hybrid without PAM and SiO\textsubscript{2} with 0.05 % PAM) were also prepared. These were then visually observed for some time. In the ultrasonication test, two steps of sonication had been employed; firstly, the dispersion of SiO\textsubscript{2} in brine after 1 h of magnetic stirring was then sonicated in ultrasonic bath for 1 h. Thereafter, it was combined with the IBS and stirred for another 12 h. The solution was then sonicated for another 1 h and allowed to stand undisturbed and sedimentation rate was studied. In the last measurements, the particle size was determined using particle size analyzer (DT 1202). The SiO\textsubscript{2}-IBS samples were placed in the measuring chamber equipped with a detector sensor to measure the particle’s Brownian motions at 25 °C, from which the size was recorded.

2.4. Foamability screening study

Approximately, 10 ml of each sample including the pure IBS (control) was transferred into the Pyrex test tube as shown in Fig. 2. The solutions were mechanical shaken for 30 s to observe the relative foamability capacity. The maximum foam height was recorded in mm.

2.5. Thermal stability screening study

Viscosity retention capacity is used to evaluate the thermal stability of the chemical formulations. The dynamic viscosity of SiO\textsubscript{2}-IBS hybrids and IBS solution were performed according to ASTM D445 using modular compact rheometer (MCQ 300). The viscosity was analyzed at 25 and 100 °C, and viscosity retention (thermal stability) capacity was computed according to Eq. 1.

\[
\text{Thermal stability} = 100 - \left[ \frac{\text{viscosity at 25°C} - \text{viscosity at 100°C}}{\text{viscosity at 25°C}} \right] \times 100
\] (1)

2.6. Statistical analysis, modeling and optimization study

Analysis of variance (ANOVA) was used to statistically analyze the data obtained. The maximum criteria for foamability and viscosity retention (thermal stability) were set as a goal to obtain an optimum condition. The highest order polynomial with the significant terms was considered by taking cognizance of the model that is not aliased.

2.7. Particle partitioning at the foam lamella

Exactly 50 ml of the optimized SiO\textsubscript{2}-IBS was placed in the graduated cylinder and vigorously shaken for 1 min to generate maximum foam volume. After 30 min, the liquid drainage was carefully poured out from the cylinder and the volume was recorded. The refractive index calibration curve of SiO\textsubscript{2} was generated (Fig. 3) to find the concentration (m/v %) of particles in the liquid drainage. This concentration was used to obtain the mass (g) of particles in the liquid drainage.

2.8. Interfacial tension (IFT) measurement

The IFTs in oil/brine, oil/SiO\textsubscript{2} and oil/SiO\textsubscript{2}-IBS systems were measured using Interfacial Tension Analyser (IFT 700) via a pendant drop. The oil droplets as shown in Fig. 4 were stabilized in the continuous phase for 1 min, and images were captured using a high resolution camera (Newport M-R565). The IFT was analyzed by the DROPimage software according to Young-laplace equation by fitting the oil drop profile with the continuous phase (solution). Thereafter, the average IFT values were recorded.
to minimize charges due to SiO$_2$ for proper viewing at different magnification from 100-100,000X.

3. Results and discussion

3.1. Result of compatibility and stability study

Compatibility study is commonly assessed by the formation of precipitates (Han et al., 2013). Precipitates occur due to lack of chemical interaction as a result of repulsive force. The SiO$_2$ used in this study demonstrated good compatibility with IBS, because no precipitates were formed in all the SiO$_2$-IBS hybrids formulated. The compatibility was further supported by FTIR experiments; the result is presented in Fig. 5. In the figure, the FTIR spectra of the pure dried SiO$_2$ and SiO$_2$-IBS hybrid were phased for easy comparison. The peaks absorbed at 1085.11 (sharped), 796.8, 563.8 cm$^{-1}$ in the FTIR of the dried SiO$_2$ were due to the Si-O-Si stretching, bending and rocking vibrations, respectively, which are also in accordance to findings in (Moore et al., 2003; Ryu and Tomozawa, 2006). Similarly, the FTIR spectrum of SiO$_2$-IBS revealed the presence of a broad peak at 3319.52 cm$^{-1}$ and additional peak at 1633.7 cm$^{-1}$ due to the presence of N-H and C=O in the IBS. It can also be seen that the peak at 796.8 cm$^{-1}$ in SiO$_2$ disappeared, and a bathochromic shift (redshift) was observed from 563.8 to 595.14 cm$^{-1}$ revealing the formation of Si-C bond between the hybrid components. Similar observations were reported in SiO$_2$-polymer hybrid from the work of Zhu et al. (2014). In addition, a sharp intensity reduction of a SiO$_2$ peak at 1085.11 cm$^{-1}$ was noticed in the SiO$_2$-IBS which further supported the chemical interaction between the SiO$_2$ and IBS.

However, the stability of SiO$_2$-IBS was carefully studied by a visual test without any pH adjustment at 25 °C. The SiO$_2$-IBS had stability up to 8 h compared to the 3 h stability of pure SiO$_2$ dispersed in brine. To improve the stability of the dispersion, additional experiments were performed as described in (Ghadimi et al., 2011). pH control, viscosity enhancement and ultrasonication were selected for further stability test. In the pH control test, the normal pH of the dispersion at 25 °C was recorded to be 6.92. After the pH was adjusted to 4, 8, 10 and 12, drastic and rapid precipitation of the dispersions...
were observed in basic medium which typically indicated the destabilization of chemical interaction between the SiO$_2$ and IBS. While at pH 4 no precipitation was seen, but the dispersion had stability within 12 h. This finding is contrary to the report in (Ghadimi et al., 2015), where the authors claimed pH 10 as the optimized stability condition for nanofluid hybrid. Ghadimi et al. (2011) had explained the concept of Stoke's law in the field of nanofluid dispersions that by increasing the viscosity of base fluids would help to decrease particles speed and improve dispersion stability. Therefore, a water soluble polyacrylamide polymer (PAM) was incorporated in the SiO$_2$-IBS hybrid mixture at concentrations of 0.05, 0.1 and 0.15 % as shown in Fig. 6. The SiO$_2$-PAM without IBS was also prepared as a control. The same concentrations of SiO$_2$ and IBS were maintained in all the mixture. It was found out that addition of PAM in the SiO$_2$-IBS blend did not improve the stability, rather, the particles started to sediment immediately within 1 h and nearly sedimented at 6 h (Fig. 6b). No further changes were noticed up to 44 h of visual observations. This happened because some polymers caused particles to flocc, which make them acquire higher density than the base fluid, and easily be dragged by gravity force (Derksen, 2014). In comparing with the SiO$_2$-PAM blend, PAM had significant stabilization effect at a concentration of 0.05 %. Similar observation of the particle stabilization with PAM was reported in the work of Sharma et al. (2016), where the authors found that, SiO$_2$ blend containing surfactant and PAM induced additional flocculation of particles, which resulted to rapid particles sedimentation. Thus, PAM in our study could only improve the viscosity of SiO$_2$-IBS, but not its stability. However, in the ultrasonication experiment, it can be seen from the same figure that SiO$_2$-IBS (OPT) had been found to be stable up to 36 h with fewer particles still remained suspended after 48 h. In addition, Fig. 7 describe the rate of particles sedimentation as a function of time, it can be observed that SiO$_2$ in brine sediment faster than the SiO$_2$-IBS (OPT) hybrid, this is due to the electrical and sound vibration from the ultrasonic bath which broke down particle aggregations in the IBS and rendered them suspended in the medium (Mahbubul et al., 2017). To further support the ultrasonication stability, particle’s size was measured. The particle size result is presented in Fig. 8. It can be seen from the figure that the particles are within the normal distribution and the agglomeration will not cause further destabilization (Chang et al., 2007).

### 3.2. X-Ray diffraction (XRD) result

The XRD result of pure dried SiO$_2$ is presented in Fig. 9a, from the figure it can be seen that SiO$_2$ exhibited characteristic of an amorphous nature due to the appearance of one broad diffraction shoulder between 20-30° at 2 theta, which is typically the characteristic of amorphous silica. However, there was a slight transformation of nature from amorphous to crystalline like nature due the appearance of crystalline peaks as indicated in the XRD spectra of the hybrid material (Fig. 9b). This transformation was due to the interaction of SiO$_2$ and IBS.

![Fig. 6: Particles stabilization by ultrasonic bath after (a) 0 h, (b) 10 h, (c) 20 h, (d) 32 h and (e) 44 h](image)

![Fig. 7: Sedimentation rate of optimized hybrid surfactant (SiO$_2$-IBS) in comparison with SiO$_2$](image)

![Fig. 8: Particle size distributions in nanofluid hybrid](image)

### 3.3. Field emission scanning electron micrograph (FESEM) result

Fig. 10 is a field emission scanning micrograph (FESEM) of pure dried SiO$_2$ nanoparticle and SiO$_2$-IBS hybrid. From the Fig. 10a and 10b, it can be seen...
that SiO$_2$ shows the morphology of network texture with pores. However, from the Fig. 10c and d of SiO$_2$-IBS, the surface network structure morphology of SiO$_2$ transformed to fine texture like and the pores were not seen, these indicated that IBS has been embedded within the structure.

![Fig. 9: XRD spectra of (a) pure dried SiO$_2$ (b) SiO$_2$-IBS hybrid](image)

**3.4. IFT result**

The IFT for oil/brine system was found to be 120.3 ± 9.8 mN/m, this represented the existing interfacial force between the oil and brine. The dispersion of SiO$_2$ in the brine significantly lowered the IFT to 31.6 ± 3.9 mN/m. This is because SiO$_2$ has the ability to orient its hydrophilic and hydrophobic like parts in the aqueous and oil phase, respectively, and induced interfacial effect (Li et al., 2013). The SiO$_2$-IBS blend had additional IFT reduction effect, and the result was found to be 10.6 ± 6.8 mN/m. This further reduction was a result of IBS surfactant in the blend, because surfactant is known as IFT reduction agents (Djemiat et al., 2015).

**3.5. Result of foamability**

The foamability data obtained were statistically analyzed and presented in Table 4. From the table, the value of prob. > F was found to be less than 0.05, indicating that the model terms (SiO$_2$, IBS, SiO$_2$*IBS, SiO$_2$* and aging) are significant. Similarly, the F value of 11.70 described the significance of the model chosen (cubic model). The R square value is used to evaluate the fitness of the model, so long it approaches unity (1.00) (Nguyen et al., 2014; Adamu et al., 2017). It can be seen that the R square of 0.8068 revealed that the model fits with the real data and this is confirmed by observing the small difference between the predicted (0.7379) and adjusted (0.5121) R square values.

![Table 4: Analysis of variance (ANOVA) for foamability](image)

The plot of residual data is shown in Fig. 11, the data can be seen to appear within the normal probability, and this further confirmed the fitness of the data obtained with the lack of fit value that is not significant. The adequate precision, which describe signal to noise ratio was found to be 12.767, and this is greater than 4 indicating the adequate signal precision. The cubic model developed is presented in Eq. 4, this describes a quantitative relationship between the foamability and the synergistic effects of the model terms (SiO$_2$, IBS, SiO$_2$*IBS, SiO$_2$ and aging). Cubic model Eq. 3;

Foamability = +89.54 +5.36[IBS] - 9.75[IBS]*[SiO$_2$] + 4.26[SiO$_2$]*[Aging] + 5.25 [IBS]*[SiO$_2$]*[Aging] (3)

Similarly, Fig. 12 shows the graphical interactions in three dimensional plots. From the graphs, it can be seen that the synergistic interaction of SiO$_2$ with IBS and aging showed that as the concentration of SiO$_2$ and IBS increases, the relative foamability of SiO$_2$-IBS hybrid also increases. This indicates the surface activity of SiO$_2$ as a result of interaction with IBS which resulted to the increase in the relative foamability of the hybrid. Vatanparast et al. (2017) also observed similar phenomenon when they studied the synergistic interaction between SiO2 and SDS surfactant. Similarly, Mo et al. (2012) established that the foam height is directly proportional to the SiO$_2$ concentrations.

**3.6. Result of particle partitioning at the foam lamella**

The result of particle partitioning at the foam lamella demonstrated that foam generated in 1 min could result to 44 % of particles to be partitioned at
the foam lamella. This indicates that the SiO$_2$ has greater effect to be adsorbed at the foam lamella and stabilize the foam.

$$\text{Thermal stability} = +15.74 + 4.56[IBS]^2 + 6.63 [IBS][SiO}_2 - 5.12[Aging] -3.38 [IBS][SiO}_2[Aging]$$

(4)

**3.7. Result of thermal stability**

Thermal stability data obtained were also statistically analyzed similar to foamability. The ANOVA result is presented in Table 5. The model (cubic) and the model terms (SiO$_2$-IBS, IBS$^2$, aging, SiO$_2$*IBS*aging) are all significant. The data were well fitted in the model and within the normal probability (Fig. 13). The synergistic effects between the model terms with respect to thermal stability were established in the developed model (Eq. 5). Fig. 14a shows that at relatively higher concentrations (#1.00) of IBS and SiO$_2$, the SiO$_2$-IBS hybrid had a maximum thermal stability at 100 °C. However, with respect to aging, the result shows that the hybrid is more stable at the early days (6 days). This indicates that aging is a crucial input parameter on the stability of hybrid. Cubic model Eq. 4:
3.8. Optimization and point prediction

After the establishment of the responses (foamability and thermal stability), the optimum condition could be achieved. The concentrations of SiO$_2$ and IBS and the aging time were kept within the range taken (-1.68 to +1.68), while the maximum value for foamability and thermal stability were set as criteria goals. Out of 18 optimum conditions generated, Table 5 shows the selected optimum condition based on the desirability which describes the extent to which goal criteria chosen to achieve the optimal conditions (Fig. 15).

To evaluate the accuracy of the developed models, an experiment was further conducted using the optimized conditions, and the result is presented in Table 6. From the table, the experimental values obtained for foamability and thermal stability were in good agreement with the values obtained from the predicted models. The minimum error observed indicates the accuracy of the cubic model for the foamability and thermal stability, respectively.

| Model | Sum of squares | Degree of freedom | Mean square | R square | Adjusted R square | Predicted R square | Adequate precision | F value | P value | Prob. > F |
|-------|----------------|--------------------|-------------|----------|-------------------|-------------------|-------------------|---------|---------|-----------|
| Model | 1105.38        | 4                  | 276.35      | 0.7781   | 0.7190            | 0.5014            | 12.343            | 13.15   | 0.001   | significant |
| A$^2$  | 305.27         | 1                  | 305.27      | 0.50      | 0.50              | 0.50              | 14.53             | 0.0017  | 0.0001  | significant |
| AB    | 351.13         | 1                  | 351.13      | 0.50      | 0.50              | 0.50              | 16.71             | 0.0009  | 0.0010  | significant |
| C     | 357.86         | 1                  | 357.86      | 0.50      | 0.50              | 0.50              | 17.03             | 0.0009  | 0.0009  | significant |
| ABC   | 91.13          | 1                  | 91.13       | 0.50      | 0.50              | 0.50              | 4.34              | 0.0548  | 0.0010  | significant |
| Residual | 315.17     | 15                 | 21.01       | 0.50      | 0.50              | 0.50              | 4.20              | 0.0632  | 0.0010  | not significant |
| Lack of fit | 281.67  | 10                 | 28.17       | 0.50      | 0.50              | 0.50              | 4.20              | 0.0632  | 0.0010  | not significant |

Table 5: Analysis of variance (ANOVA) for thermal stability

A = [IBS], B = [SiO2] and C = Aging

| Input variables (coded values) | Desirability | Predicted values | Experimental values | Error (%) |
|-------------------------------|--------------|------------------|---------------------|----------|
| IBS % | SiO2 % | Aging (days) | Foamability (mm) | Thermal stability (%) | Foamability (mm) | Thermal stability (%) |          |
| 1.0  | 1.0   | 1.0            | 0.681              | 96.4                | 23.1     | 100                | 25.4          | 3.6  | 9.1 |

Table 6: Optimized conditions of SiO$_2$-IBS hybrid

- This novel hybrid material could find wide applications in the EOR, particularly, in the higher temperature reservoirs.

Fig. 13: Normal residual plot of thermal stability data

4. Conclusion

The following conclusions can be deduced from this study:

- A new EOR hybrid formulation (SiO$_2$-IBS) had been successfully developed, optimized and characterized.
- The effect of SiO$_2$ nanoparticles in the hybrid had been established to improve the relative foamability, thermal stability and IFT reduction between crude oil and brine.
- Two multivariate models (cubic) were developed and validated, and the synergistic interaction between the components were quantified.
Fig. 14: The three dimensional graphical plots of thermal stability showing the synergistic interaction of SiO$_2$ with IBS (a), SiO$_2$ with aging (b) and IBS with aging (c).

Fig. 15: Desirability response for optimum condition

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