

Original Research Article

Performance Assessment of Biomass-derived Urea-Furfuraldehyde Resins as Oilfield Scale Inhibitors

Abstract

Oilfield-scale formation is a persistent challenge in the oil industry. While numerous scale inhibitors have been in use for decades, there is a significant research gap in discovering renewable, cost-effective, ecologically friendly, and efficient inhibitors. This study sets out to fill this gap by investigating the potential of scale inhibitors (SIs) made from bio-resin derivatives of red onion skin; ROF and ROFU, in reducing CaCO_3 and CaSO_4 scales. The scale inhibition performance of ROF and ROFU was rigorously evaluated using the NACE standard static bottle test method. The data revealed that increasing scale-inhibitor contact time, concentration, and temperature enhances inhibitor efficacy, with the best inhibition efficiencies found for ROF and ROFU on the CaSO_4 scale compared to the CaCO_3 scale across all studied parameters. A comparison of the prepared SIs with a commercial scale inhibitor (CSI) showed a high inhibition rate on all scales tested. Despite having a lower inhibition rate (IE) than CSI, ROF and ROFU demonstrate significant potential as green oilfield SIs. This sustainable technique, which transforms agro-waste into a valuable oilfield chemical via a one-pot chemical process, could have profound economic and societal benefits, offering hope for a more sustainable future in the oil industry.

1. Introduction

A sustainable future may see a large contribution from biomass, an adaptable and renewable resource. Agricultural food wastes and by-products with significant bioactive chemicals are produced in large quantities worldwide, particularly along the whole supply chain. Evaluating food waste as valuable biomass that may be turned into profitable products rather than an unmanageable waste stream has begun (Hatti-Kaul *et al.*, 2007; Ituen *et al.*, 2017). Food waste can produce biofuels, enzymes, antioxidant extracts, new biodegradable materials,

electricity, and other commercial items because it is cheap and renewable (Verma *et al.*, 2024).

Biobased chemicals are drawing more attention as sustainable substitutes in several industries, including oil and gas (Benyus, 2001). Within the petroleum sector, oilfield scale deposition is a frequent difficulty that results in lower production efficiency and higher operating expenses. Scale formation happens when precipitated minerals, like calcium carbonate (CaCO_3), calcium sulphate (CaSO_4) and/or barium sulfate (BaSO_4), settle on wellbore surfaces, pipelines, and downhole equipment from generated water. These deposits can harm equipment, drastically lower flow rates, and plug holes, resulting in expensive interventions. Scale development is a major concern in oilfield operations, impacting equipment integrity and production efficiency. Traditional scale inhibitors are often made from petroleum-based compounds, despite rising interest in sustainable alternatives, however, the limited availability and environmental impact of chemicals derived from fossil fuels have spurred the investigation of biobased substitutes (Powell *et al.*, 2011; Jessop *et al.*, 2015, 2018; Li *et al.*, 2019).

Green oilfield scale inhibitors made of biobased materials provide several environmental and sustainability advantages. The carbon footprint associated with oilfield activities can be greatly decreased by substituting renewable chemicals for petroleum-based ones. Additionally, biobased inhibitors provide a more sustainable method of producing oil and gas and help protect nonrenewable resources (Abdel-Raouf *et al.*, 2021; Verma & Fortunati, 2019). These environmentally friendly substances may also be biodegradable, lessening their negative effects on aquatic environments and ecosystems. In conclusion, a viable path toward environmentally responsible and sustainable operations in the oil and gas sector is using biobased compounds as green oilfield scale inhibitors as these substitutes have the ability to lessen scale deposition while also lessening the environmental effect of conventional petroleum-based inhibitors by utilizing renewable resources (Dickinson *et al.*, 2011).

Red onion skin (ROS) is one of the renewable feedstocks that has been used to create biomass-derived resins, such as ROSE-glutaraldehyde (ROG), ROSE-furfuraldehyde (ROF), and ROSE-urea furfuraldehyde (ROFU) resins (Victor-Oji *et al.*, 2024; Obuebite *et al.*, 2023a & 2023b, Phung Hai *et al.*, 2021; Tenorio-Alfonso *et al.*, 2020). In ROFU resins, furfuraldehyde and urea react via a one-pot, two-step process to generate a crosslinked polymer structure, which has the potential to mitigate scale formation due to their distinct chemical makeup. Their regenerative nature and possible scale inhibition features make them an attractive alternative (Husna *et al.*, 2022). By comparing ROFU resins to their conventional counterpart using the National Association of Corrosion Engineers (NACE) standard bottle tests under simulated oilfield conditions, this work explores the possibilities of bio-based resins generated from urea-furfuraldehyde as oilfield scale inhibitors, considering their synthesis, characterization, and effectiveness in preventing scale formation in terms of its inhibition efficiency, compatibility and thermal stability in other chemicals, and likely environmental impact.

The utilization of bio-based derived resins of this kind, which have not been thoroughly investigated for this purpose, makes this study innovative, and the findings of this study will advance sustainable solutions in petroleum and gas production and contribute to creating scale inhibitors for the oil and gas sector that are biodegradable and more environmentally friendly substitutes for currently used inhibitors by lowering their environmental impact and increasing production efficiency.

2. Materials and Methods

2.1 Materials

Red onion skin (ROS) was sourced locally from the Oil Mill Market (4.8585°N, 7.0648°E) in Obio-Akpor Local Government Area, Rivers State, Nigeria. The analytical grade chemicals purchased from local suppliers include Urea, Furfuraldehyde, Sodium Hydroxide, Acetone, Sodium chloride, Calcium chloride dihydrate, Magnesium chloride hexahydrate, Sodium

sulphate, sodium hydrogen carbonate, and Sodium carbonate. They were used as received without further purification.

2.2 Methods

The methodology employed during this study includes extraction of the biomass (Red onion Skin, ROS), chemical modification of ROS extract (ROSE), characterization of the chemically modified extracts, formulation and characterization of synthetic brine, formulation of the scale inhibitor (SI) solutions, and comparative assessment of the formulated scale inhibitor with a commercial scale inhibitor via static bottle test.

2.2.1 Extraction of ROS

The inedible outer skin of the red onion was thoroughly selected, cleaned, and sun-dried for about 6 hours. The dried red onion skin (ROS) was milled, sieved to 150 μ m particle size, and extracted by solvent extraction using acetone as described by Una *et al.* (2021). The ROS extract, ROSE, was concentrated using a rotary evaporator and dried in an oven at 55 °C. The dried extract was stored in an airtight container labeled ROSE for further use.

2.2.2 Formulation of Synthetic brine

Synthetic brine was simulated in the laboratory to replicate formation brine by dissolving specific inorganic salts in distilled water according to the National Association of Corrosion Engineers (NACE) standard method, TM0197-2010. Synthetic brine was prepared by mixing solutions A and B, defined as calcium-containing brine and carbonate or sulfate-containing brine for CaCO₃ and CaSO₄ scale types, as outlined in Table 1, with distilled water in a 1:1 ratio. The formulated brines for CaCO₃ and CaSO₄ were filtered through a 0.45 μ m membrane filter paper to remove undissolved salts.

Table 1 Composition of Synthetic water and test condition

Scale Type	Brine Concentration, g/L
CaSO ₄	A. Calcium Containing Brine: NaCl = 7.50, CaCl ₂ .2H ₂ O = 11.10

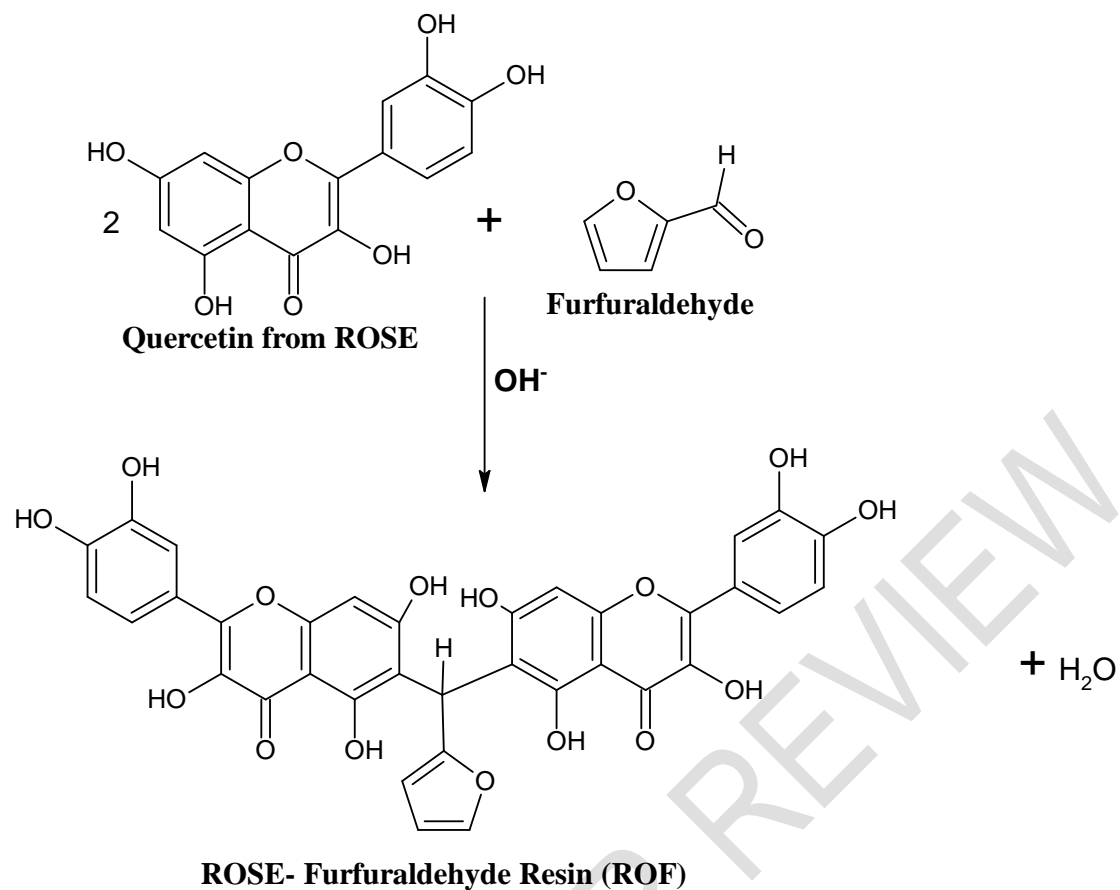
(A+B)	B. Sulphate Containing Brine: NaCl = 7.50, Na ₂ SO ₄ = 10.66
CaCO ₃	A. Calcium Containing Brine: CaCl ₂ .2H ₂ O = 12.15, MgCl ₂ .6H ₂ O = 3.68
(A+B)	B. Carbonate Containing Brine: NaHCO ₃ = 7.36, NaCO ₃ = 33.0

2.2.3 Synthetic Brine Characterization

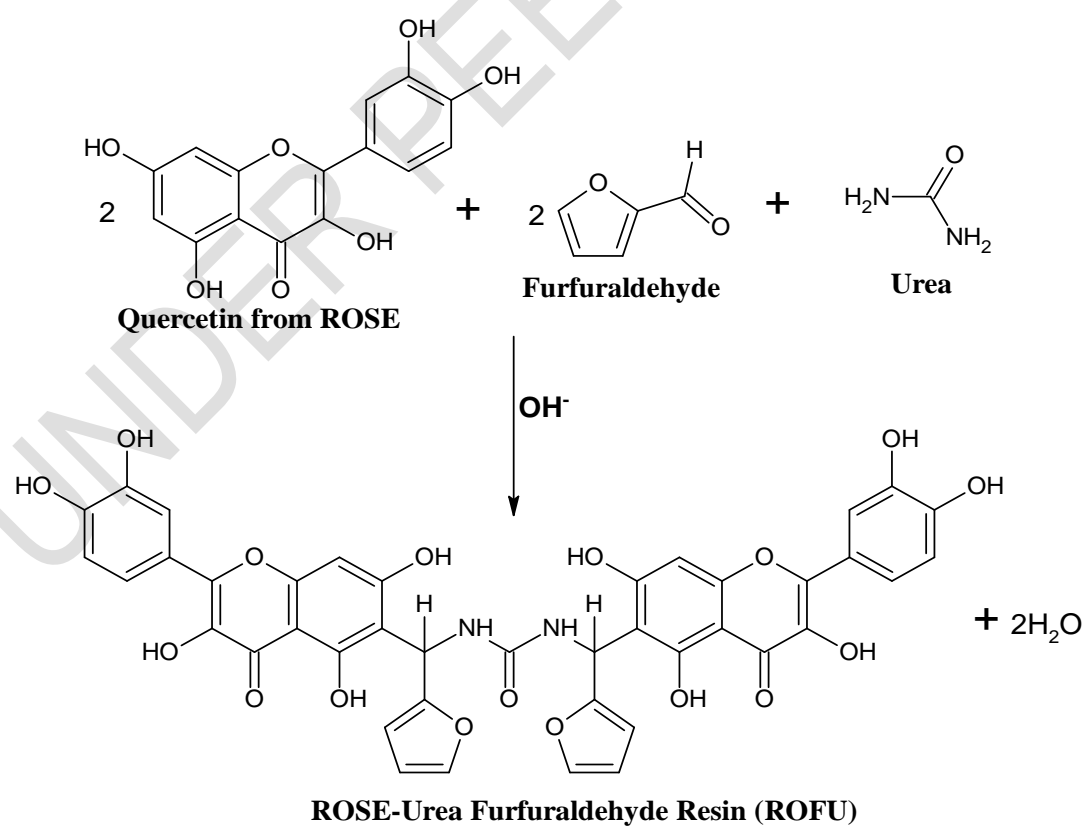
The physicochemical properties of synthetic brine, such as pH, density, salinity, conductivity, turbidity, dissolved oxygen, and alkalinity, were electrometrically analyzed for each studied scale type using the Horiba water monitor (U-52 model).

2.2.4 Chemical Modification of ROSE

The biomass extract, ROSE, was modified via a one-pot condensation reaction with varying mixing ratios of ROSE: Furfural and ROSE: Furfural: Urea in an alkaline catalyst to give ROF and ROFU, respectively. For ROF, a reaction mixture of ROSE: Furfuraldehyde (2:1 mole ratio) was charged into a pre-weighed reactor vessel gradually heated to 120°C with continuous stirring under reflux for 60 min in the presence of a catalytic quantity of NaOH (based on ROSE), while for ROFU, a reaction mixture of furfuraldehyde: urea (2:1 mole ratio) was charged into a pre-weighed reactor vessel and refluxed continuously at a temperature of 70°C for 30 mins after which 2 moles of ROSE and a catalytic amount of NaOH was introduced into the mixture. This mixture was refluxed for 60 min with continuous stirring at 70°C; at the end of the reaction (when the volume of water condensed in the dean and stark trap is constant), the products were cooled, weighed, and stored in airtight containers labeled ROSE-furfuraldehyde (ROF) and ROSE-Furfuraldehyde-Urea (ROFU) resin respectively for later use. The proposed synthetic routes for ROF and ROFU are depicted in Schemes 1 and 2.



Scheme 1 Proposed reaction of ROSE with Furfuraldehyde in 2:1 molar ratio



Scheme 2 Proposed reaction of ROSE with Furfuraldehyde and Urea in 2:2:1 molar ratio.

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2.2.5 Characterization of ROF and ROFU

ROSE, ROF, and ROFU were characterized using an Agilent spectrophotometer scanning in the 4000 – 650 cm^{-1} range to detect the presence of functional groups. The pH, color, appearance, and solubility of ROSE, ROF, and ROFU in water were also determined. Scanning Electron Microscopy (SEM) was used to determine the morphology of the scale crystals formed before and after precipitation. This characterization was done for the calcium sulfate and carbonate-containing brines in the presence and absence of the formulated scale inhibitors ROF and ROFU.

2.2.6 Scale Inhibitor Evaluation Tests

The modified ROSE derivatives, ROF and ROFU, were evaluated to ascertain their competency to function as scale inhibitors. The assay, which includes pH, thermal stability, and compatibility tests, determines whether they meet the set criteria for scale inhibitors before performance evaluation via the static bottle test method. The procedure for each test is outlined below, and the pass or fail criterion for each test is given in Table 2.

- (a) **pH tests:** The pH values were measured using an ORP pH meter calibrated with two buffer solutions of pH = 4.0 and 10.0.
- (b) **Thermal Stability tests:** Fifteen milliliters of each test inhibitor were introduced into Erlenmeyer flasks and examined for clarity. They were then transferred to a high-pressure, high-temperature (HPHT) vessel and heated to 130°C for 24 hours. After that, the vessel was taken out of the oven and placed in a desiccator to cool for two hours. The formulated inhibitors were thereafter transferred to a clear bottle and examined for changes in appearance because of the high-temperature impact on the solutions.
- (c) **Brine compatibility tests:** The brine compatibility with the formulated scale inhibitors and SIs was evaluated at 25°C and 90°C. Before the compatibility test, the synthetic brine was filtered using a 45 μ filter paper into a transparent 15 mL glass tube. The formulated SIs were then mixed with brine to cover the range of concentration studied (20 ppm - 100 ppm).

The glass tubes were checked for signs of incompatibility, such as cloudiness/haziness and precipitation.

Table 2: Inhibitor Evaluation Criterion

TEST	PASS	FAIL
pH for 100% product	> 5	< 5
Thermal stability at 130 °C	Clear with or without a change in the color	Precipitation
Compatibility with water	Clear solution for all ranges of concentration (20ppm - 100 ppm) at both 25 °C and 90 °C	Slightly Hazy, Hazy, or precipitated solution

2.2.7 Scale Inhibitor Performance Evaluation Test

Standard test methods were used to evaluate the scale inhibition potential of the formulated scale inhibitors to mitigate the precipitation of oilfield scales under laboratory conditions. This study utilizes the National Association of Corrosion Engineers (NACE) standard static bottle test method, NACE TM0374-2007, to assess the inhibitor's ability to mitigate precipitation of calcium sulfate and calcium carbonate scales from solution at various scale inhibitor concentrations under static laboratory conditions as well as their thermal stability under the conditions listed in Table 3. The static bottle test involves incubating scaling brines mixed in 100 ml Erlenmeyer bottles for a stipulated period before being dosed with the formulated inhibitors, as shown in Figure 1.



Figure 1 Static bottle testing

The precipitation of CaCO_3 and CaSO_4 salts from the brine, if any, is followed by a measurement of the concentration of calcium ions as a function of time as described in

American Society for Testing and Materials (ASTM) test methods D511. The average Ca^{2+} values in mg/L of CaSO_4 or CaCO_3 retained in solution for the blank and each studied test concentration is defined as the percentage inhibition efficiency (IE) of the inhibitor and calculated using Equation 1:

$$\% \text{ Inhibition} = \frac{C_a - C_b}{C_c - C_b} \times 100 \quad \mathbf{1}$$

Where: $C_a = \text{Ca}^{2+}$ concentration in the treated sample after precipitation

$C_b = \text{Ca}^{2+}$ concentration in the blank after precipitation

$C_c = \text{Ca}^{2+}$ concentration in the blank before precipitation

Table 3: List of evaluated products and evaluation parameters

Product	ID	Evaluated Scale types	Evaluation conditions
ROSE-Furfuraldehyde resin	ROF	CaCO_3 and CaSO_4 scales	71 °C & 90 °C, 22 hrs
ROSE-Urea -Furfuraldehyde resin	ROFU		71 °C, 4 hrs
Commercial scale inhibitor	CSI		

3. Results and Discussion

3.1 Synthetic Brine Characterization

The physicochemical analysis of laboratory-simulated produced water and seawater was done, and the results are presented in Table 4.

Table 4 Physico-chemical assay of the synthetic brine

Parameter	Scale Type	
	CaSO_4	CaCO_3
pH	8.27	7.69
Density (g/cm^3)	1.026	1.036
Temp °C	25.3	25.1
Salinity	11.6	29.8
Turbidity (NTU)	171	185
Conductivity (ms/cm)	19.5	45.8
DO (mg/l)	7.72	7.76
Total Alkalinity (mg/l)	19	0.6

3.2 Characterization of ROF and ROFU

The pH, color, appearance, and solubility of ROSE, ROF, and ROFU in water have been assessed. The results show they are very soluble in water, dark brown in color, and have pH values of 8.0, 7.5, and 7.5, respectively, higher than the norm for scale inhibitors (pH > 5) (Table 2). Hence, they are ideal for use as oilfield scale inhibitors.

The bio-derived resins were subjected to FTIR spectroscopic analysis to confirm the presence of appropriate functional groups. The FTIR spectra of furfuraldehyde, urea, ROSE, and the resins, ROSE-furfuraldehyde resin (ROF) and ROSE-furfuraldehyde-urea resin (ROFU), are shown in Figures 2 and 3, respectively. For ROF, a broad absorption peak due to phenolic O–H stretch vibration was observed at 3272 cm^{-1} , followed by strong doublet absorption peaks at 2851 cm^{-1} and 2918 cm^{-1} due to aromatic C–H and C=O stretch vibrations, which are characteristic of quercetin structure. A mix of aryl conjugated C=C and C=O stretch vibrations were observed at 1622 cm^{-1} and 1562 cm^{-1} alongside medium absorption bands at 1462 cm^{-1} and 1443 cm^{-1} , which match aromatic C=C and =C–H stretch vibrations, the bands observed at 1380 cm^{-1} and 1339 cm^{-1} correspond to a combination of aryl O–H deformation and C–O stretch vibrations. Aromatic =C–H in-plane deformation vibration and out-of-plane C–H deformation within the furan ring were observed at 1272 cm^{-1} and 1175 cm^{-1} , respectively, confirming the bond formation between ROSE and furfuraldehyde. The weak absorption band at 1052 cm^{-1} is due to ether C–O stretch vibration, while that at 1019 cm^{-1} corresponds to the furan ring's =C–H stretch vibrations. Strong to weak absorption bands seen at 888 cm^{-1} and 784 cm^{-1} correspond to aromatic and out-of-plane C–H stretching and deformation (due to isolated H atoms) vibrations, respectively, while the band at 724 cm^{-1} is due to O–H out-of-plane bending vibrations. It is important to note that the disappearance of the characteristic C=O stretch vibrations of the aldehydic group in furfuraldehyde at 1782 cm^{-1} and 1670 cm^{-1} further affirms bond formation between ROSE and furfuraldehyde (Obuebite *et al.*, 2023b).

For ROFU, the overlying broad absorption peaks at 3350 cm^{-1} , 3290 cm^{-1} , and 3134 cm^{-1} coincide with N–H and phenolic O–H stretch vibrations in the ROSE-furfuraldehyde-urea resin structure (Scheme 2). The quercetin-characteristic doublet absorption peaks observed at 2851 cm^{-1} and 2918 cm^{-1} conform to aromatic C–H and C=O stretch vibrations, respectively. A mix of aryl conjugated $\text{C}=\text{C}$ and $\text{C}=\text{O}$ stretch vibrations was observed at 1622 cm^{-1} and 1562 cm^{-1} ; however, the band at 1592 cm^{-1} corresponds to N–H deformation and C–N stretch vibration, while the appearance of a sharp peak at 1465 cm^{-1} corresponds to the asymmetric N–CO–N stretch vibration, confirming the occurrence of an amide bond within the ROFU resin structure.

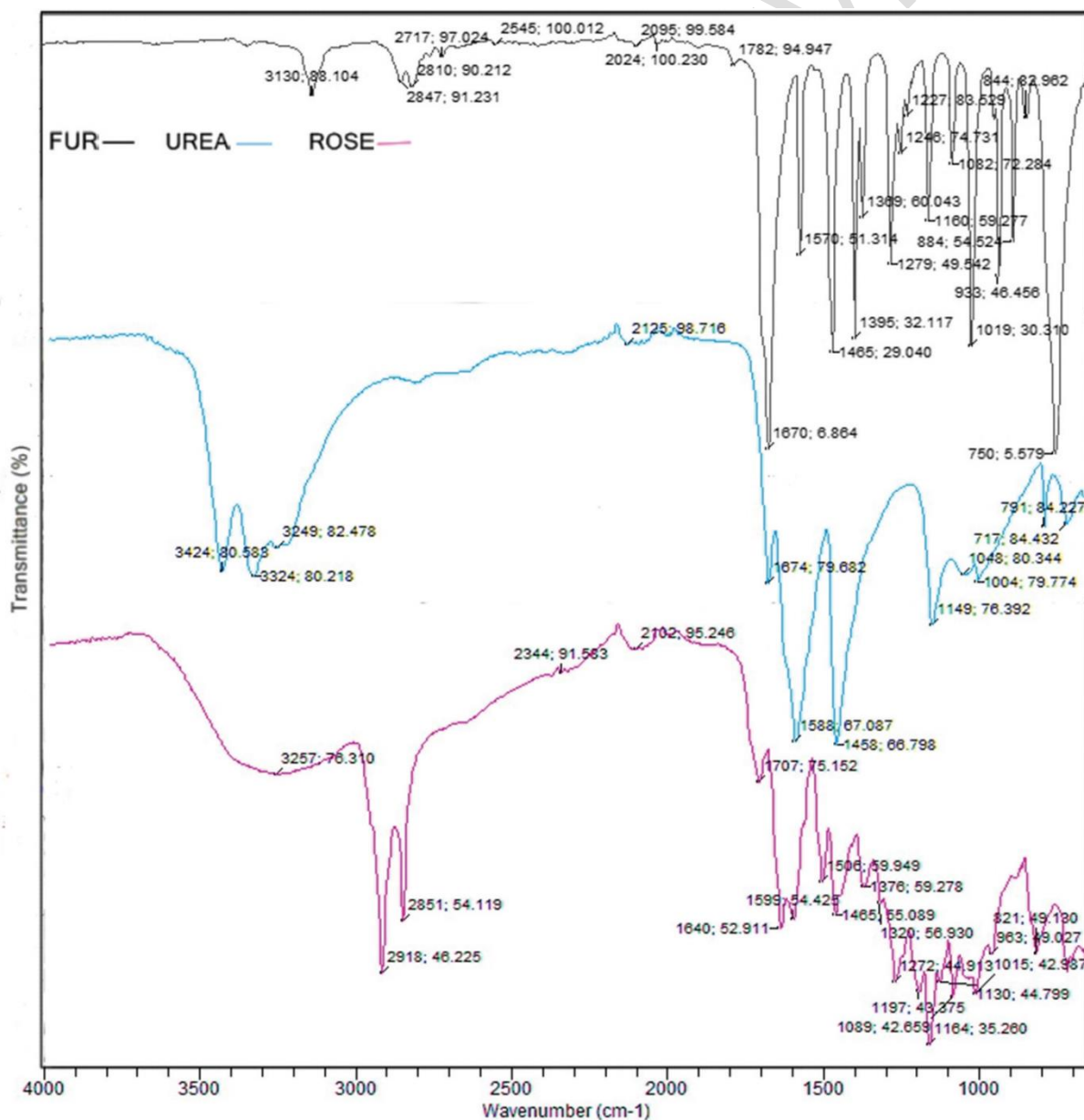


Figure 2 FTIR Spectra of Furfuraldehyde (FUR), Urea, and Red Onion Skin Extract (ROSE)

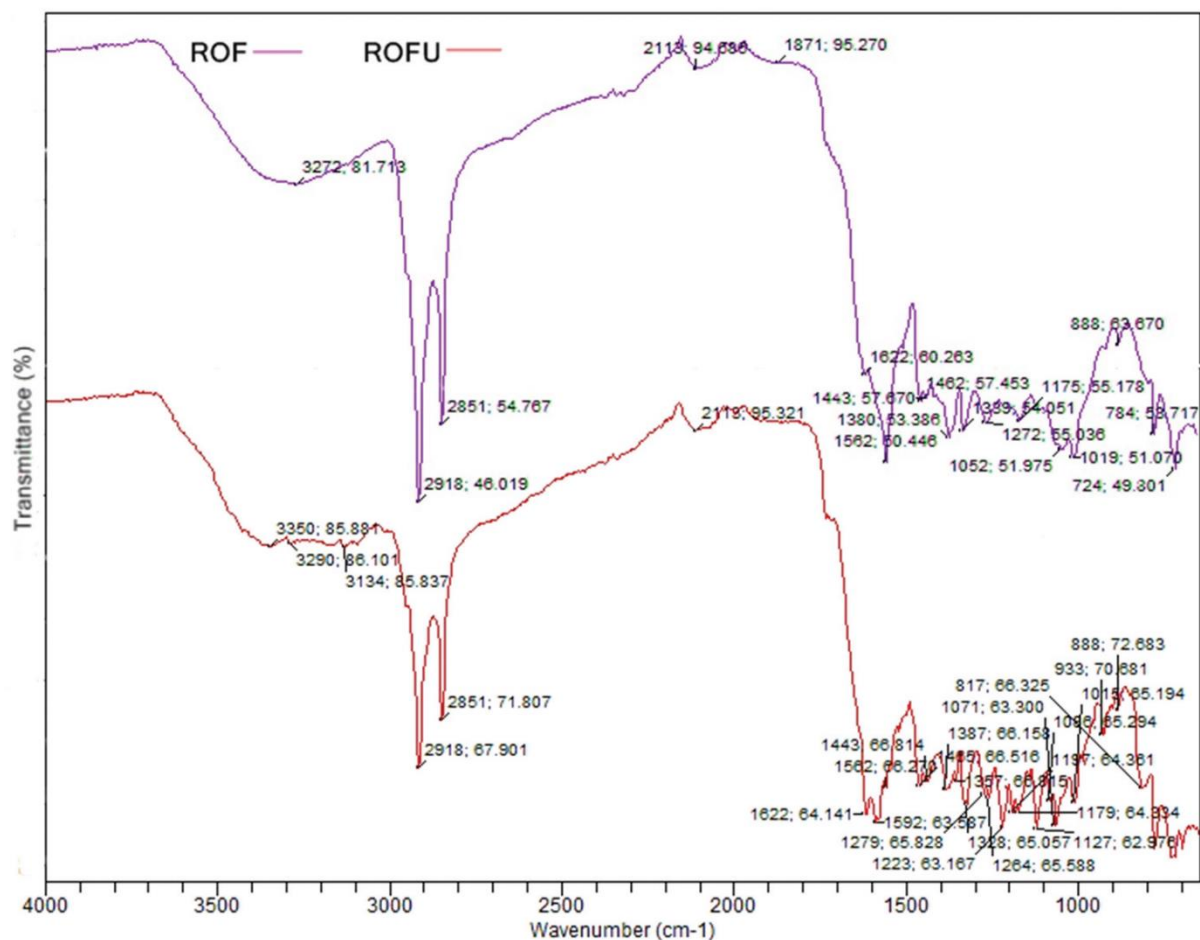


Figure 3 FTIR spectra of ROSE-furfuraldehyde resin (ROF) and ROSE-formaldehyde-urea resin (ROFU)

Stretch vibrations of aromatic C=C and =C-H in the furan ring were observed at 1443 cm⁻¹, while aryl O-H deformation and C-O stretch vibrations for the flavone ring occurred at 1387 cm⁻¹. Medium doublet peaks at 1357 cm⁻¹ and 1328 cm⁻¹ occur due to C-N stretch vibrations, while aromatic =C-H in-plane deformation vibration was observed at 1279 cm⁻¹. Weak absorption bands due to ether C-O stretch vibrations in the furan and flavone rings were also observed at 1264 cm⁻¹, 1223 cm⁻¹, 1197 cm⁻¹, 1179 cm⁻¹, 1127 cm⁻¹, and 1096 cm⁻¹, respectively. Vibrations due to methine C-H deformation and aromatic =C-H stretch in the furan and flavone rings were observed at 1071 cm⁻¹ and 1015 cm⁻¹, respectively. The absorption peaks at 933 cm⁻¹ and 888 cm⁻¹ conform to out-of-plane aromatic C-H

deformation vibrations (due to isolated H atoms) around the rings, while that at 817 cm^{-1} corresponds to ether C–O–C stretching vibrations (Obuebite *et al.*, 2023b).

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3.3 *Scale Inhibitor Evaluation*

The thermal stability and compatibility of the formulated scale inhibitors (SIs); ROF and ROFU, in the synthetic brine, were evaluated under laboratory conditions. They were found to be thermally stable after thermal exposure to temperatures up to 130°C, as no precipitation was observed, thus passing the criterion as depicted in Table 2. The compatibility tests with the synthetic brine were conducted at 25 °C and 90 °C for 24 hours and 15 minutes, respectively. No precipitation was observed for the studied period and specific temperatures, as the inhibitor solutions were clear, which signifies brine compatibility.

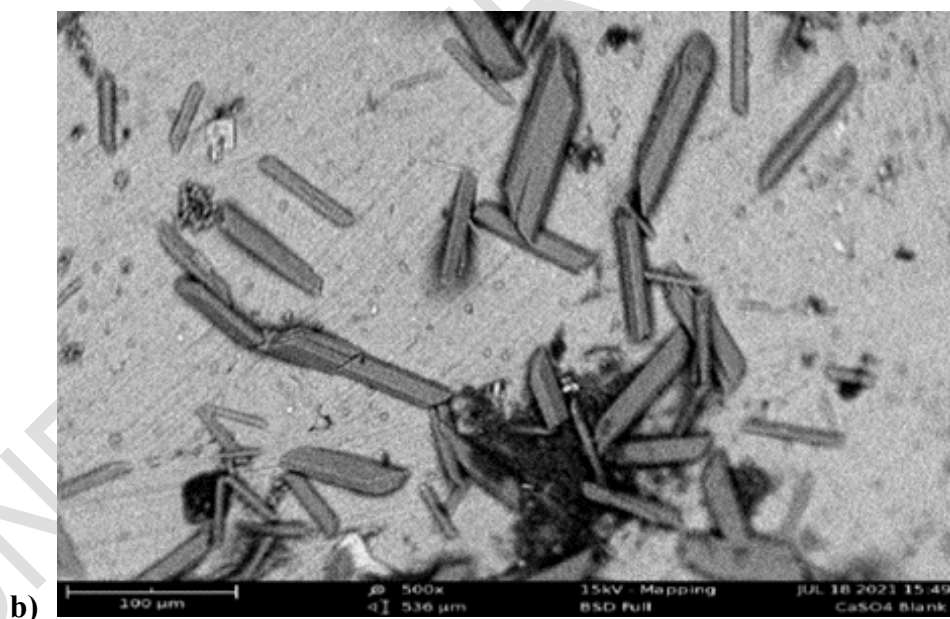
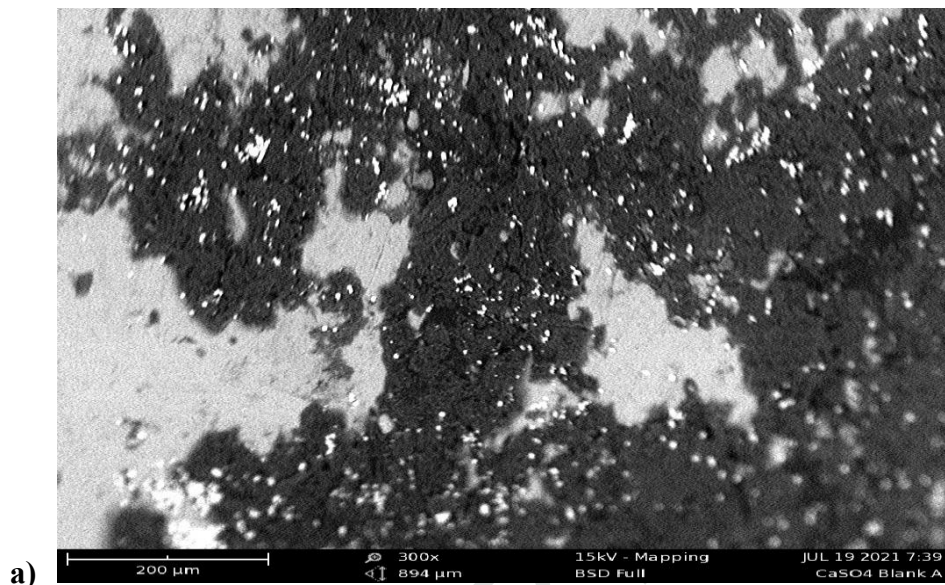
3.4 *SEM Micrographs of Scale Inhibition with ROF and ROFU*

When a scale inhibitor is introduced to a sample brine, the size and shape of the scale crystal can indicate the degree of inhibitor dispersion. As posited by Mavredaki et al. (2011), the higher the degree of crystal distortion, the smaller the crystal size, the greater the dispersibility, and the better the scale inhibition performance.

The SEM micrographs of CaSO₄ and CaCO₃ scales before and after conditioning in the absence and presence of low dosage of the prepared inhibitors, ROF and ROFU, are illustrated in Plates 1 and 2. The loose-scale crystals in the incubated blank (Plates 1a and b) became denser with well-defined crystal morphology on the addition of ROF to the CaSO₄ brine (Plate 1c). However, with the addition of ROFU, the scale crystals are decreased in size and density and are well dispersed (Plate 1d). They are also rhomboid-like, which indicates scale crystal growth inhibition (Dickinson *et al.*, 2011; Shen *et al.*, 2012).

Unlike the CaSO₄ brine, the scale crystals in the CaCO₃ brine are loosely but densely packed before incubation (Plate 2a); however, after incubation, they become dense clusters with a defined shape (Plate 2b). The introduction of ROF decreases the density of scale crystals, causing them to appear more rounded and well-dispersed with a smooth texture (Plate 2c). In contrast, with the addition of ROFU, a more dispersed and uniform distribution of the crystals is observed as well as decreased crystal density and finely distributed aggregates in smaller clusters implying suppressed crystal nucleation and growth.

From the SEM images in Plates 1 and 2 for ROF and ROFU, it may suggest that their chemical structures impacted the growth of the scale crystals as their addition to the brine either enhanced/reduced the diameter of CaCO_3 and CaSO_4 scale crystals as applicable (El-Said *et al.*, 2009; Ohimor *et al.*, 2019; Hoang, 2022).



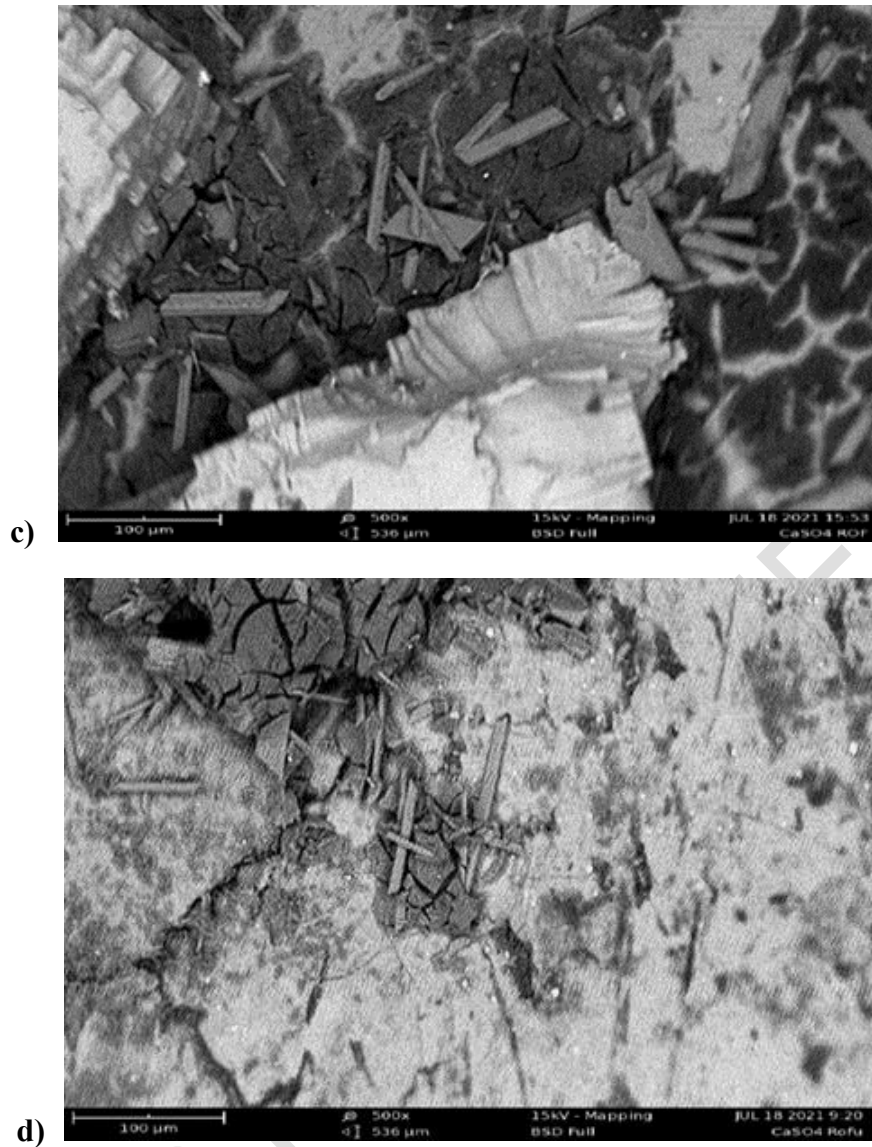
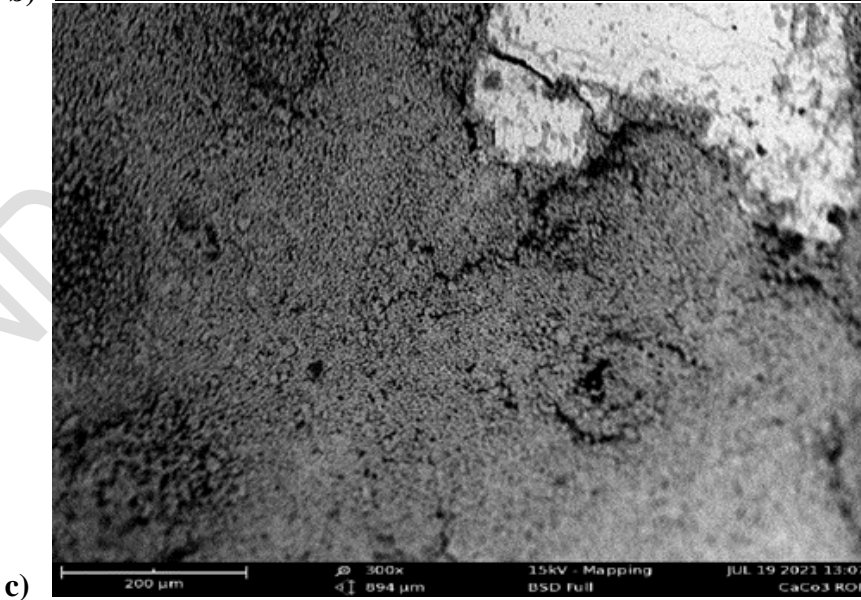
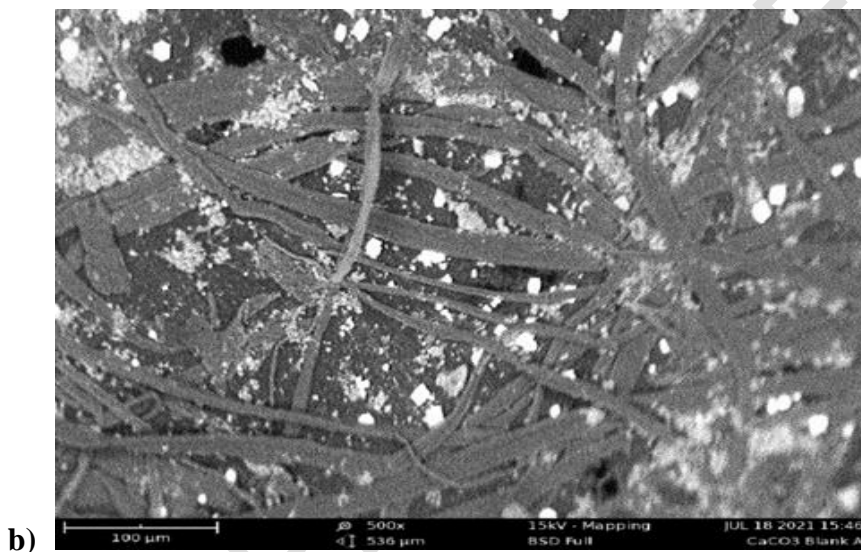
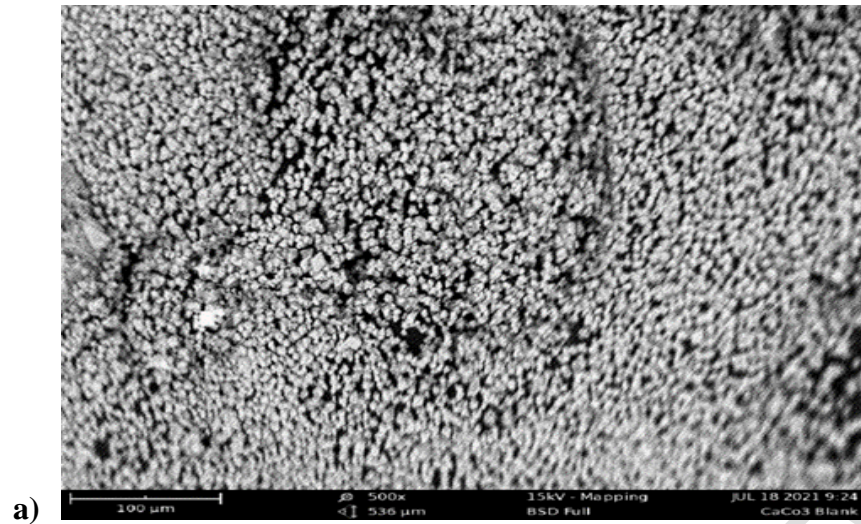


Plate 1 SEM micrograph for CaSO₄ scale deposit (a) before conditioning, (b) after conditioning, (c) inhibited with ROF, and (d) inhibited with ROFU.



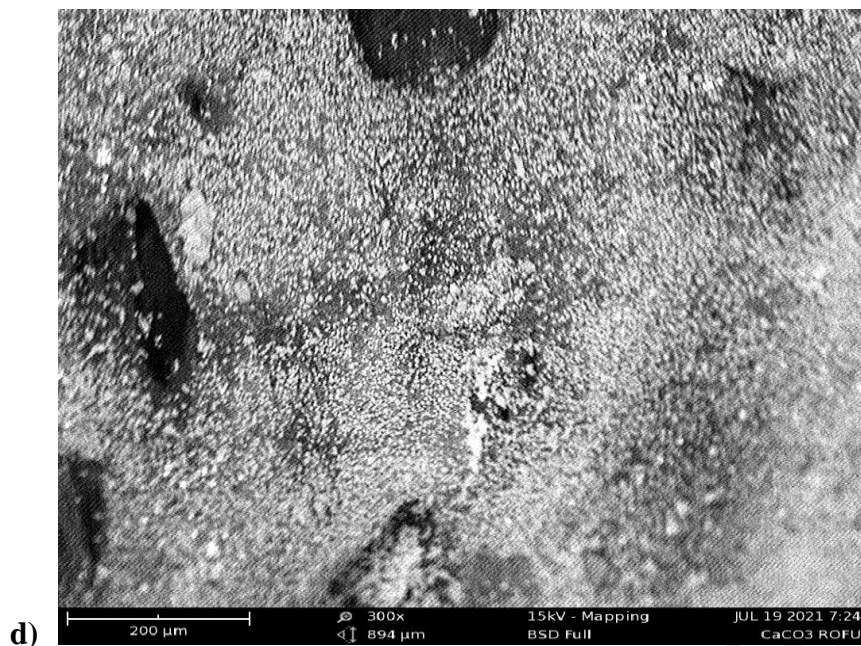


Plate 2 SEM micrograph for CaCO₃ scale deposit (a) before conditioning, (b) after conditioning (c) inhibited with ROF, and (d) inhibited with ROFU.

3.5 Performance Evaluation of ROF and ROFU as Scale Inhibitors

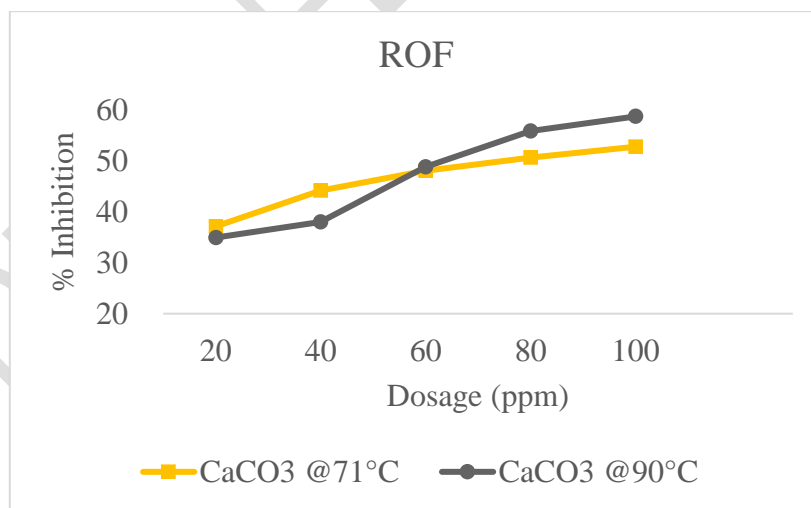
The inhibition performance efficiency (IE) of the formulated inhibitors, ROF and ROFU, was evaluated compared to a commercial scale inhibitor (CSI). The data plots in Figures 4 and 5 reveal the effect of temperature (71°C and 90°C). The data plots in Figures 4 to 9 contact time (4 and 22 hours at 71°C) and inhibitor concentration (20 to 100 ppm) on the performance of ROF, ROFU, and CSI in inhibiting CaSO₄ and CaCO₃ scales.

3.5.1 Temperature effect

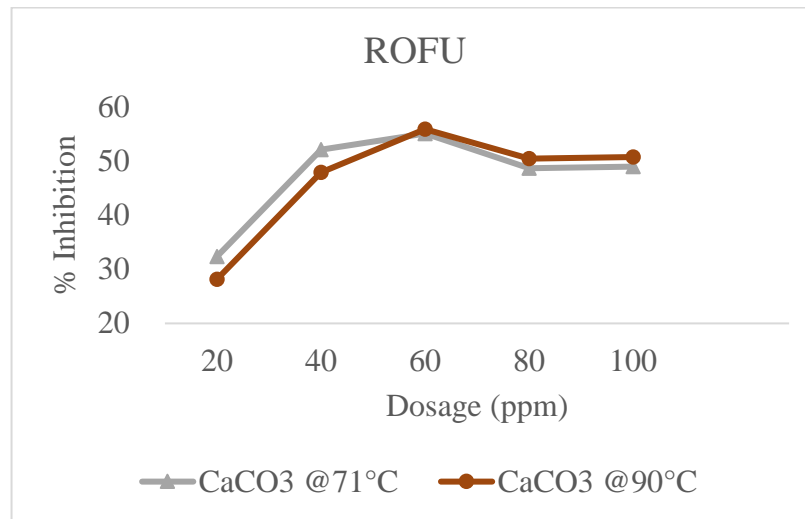
Figures 4 and 5 show comparative plots illustrating the temperature effect of ROF, ROFU, and CSI inhibition performance at 71 °C to 90 °C for CaCO₃ and Figure 5 for CaSO₄ scales, respectively. For ROF, a slight decrease in CaSO₄ scale inhibition values of 69.15 % and 68.69 % at 100 ppm and 80 ppm was observed at 71 °C and 90 °C, respectively (Figure 5a). This reflects a negligible drop with an increase in temperature. However, an increase in scale inhibition performance from 52.86 % to 58.82 % at 100 ppm was observed for CaCO₃ scales as the evaluation temperature increased from 71 °C to 90 °C (Figure 4a). This aligns with studies that indicate elevated temperatures enhance the growth of CaCO₃ scales, thus

requiring increased inhibitor concentration for better efficiency (Dyer & Graham, 2002; Olajire, 2015; Zhao *et al.*, 2019; Victor-Oji *et al.*, 2024).

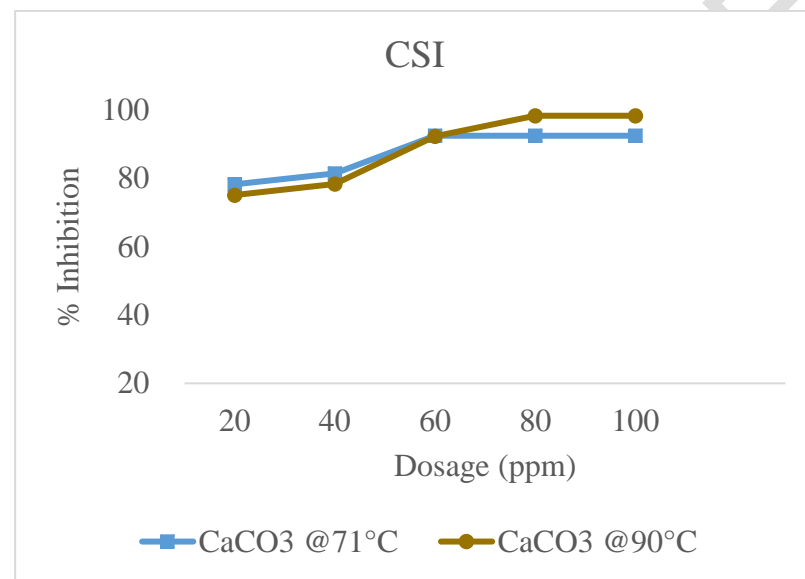
A similar trend was also observed for ROFU and CSI, as their inhibition performance in CaCO_3 scales increased from 48.92% and 92.50 % (at 71 °C) to 50.73% and 98.36% (at 90 °C) (Figure 4b and 4c) and also decreased in CaSO_4 scales from 60.38% and 75.26% (at 71 °C) to 55.07% and 68.51% (at 90 °C), respectively at 100 ppm (Figure 5b and 5c). The evaluated scale inhibitors, ROF, ROFU, and CSI, showed optimal inhibition values in CaSO_4 scales at 71 °C, while in CaCO_3 scales, optimal inhibition values were observed at 90 °C. Summarily, optimal inhibition rates were observed at higher temperatures of 90 °C than at 71 °C, suggesting that temperature increase influenced the scale inhibitor's performance, causing more inhibitors to be adsorbed onto the scale crystal. Compared to the higher inhibition values of CSI, we can infer that the inhibition performance of ROF and ROFU is above average and favorably inhibited CaSO_4 scales than CaCO_3 scales.



a)

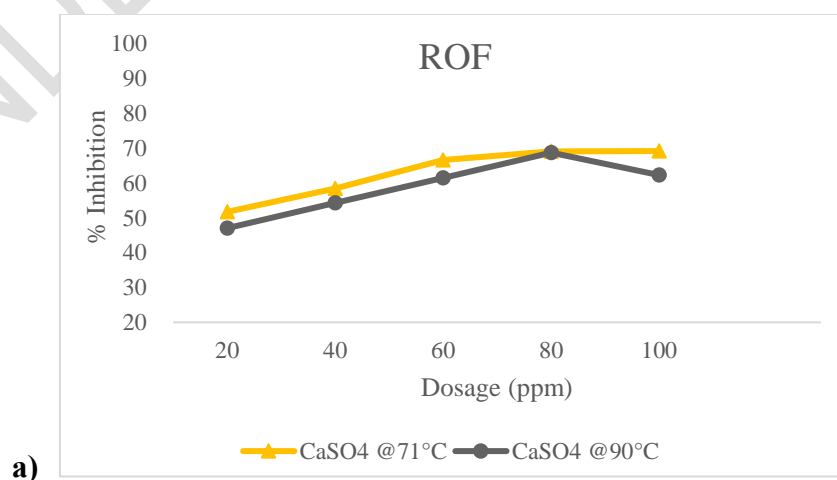


b)



c)

Figure 4 Effect of temperature on the performance of (a) ROF, (b) ROFU, and (c) CSI on CaCO₃ scales at 71°C and 90°C at 22 hrs, respectively



a)

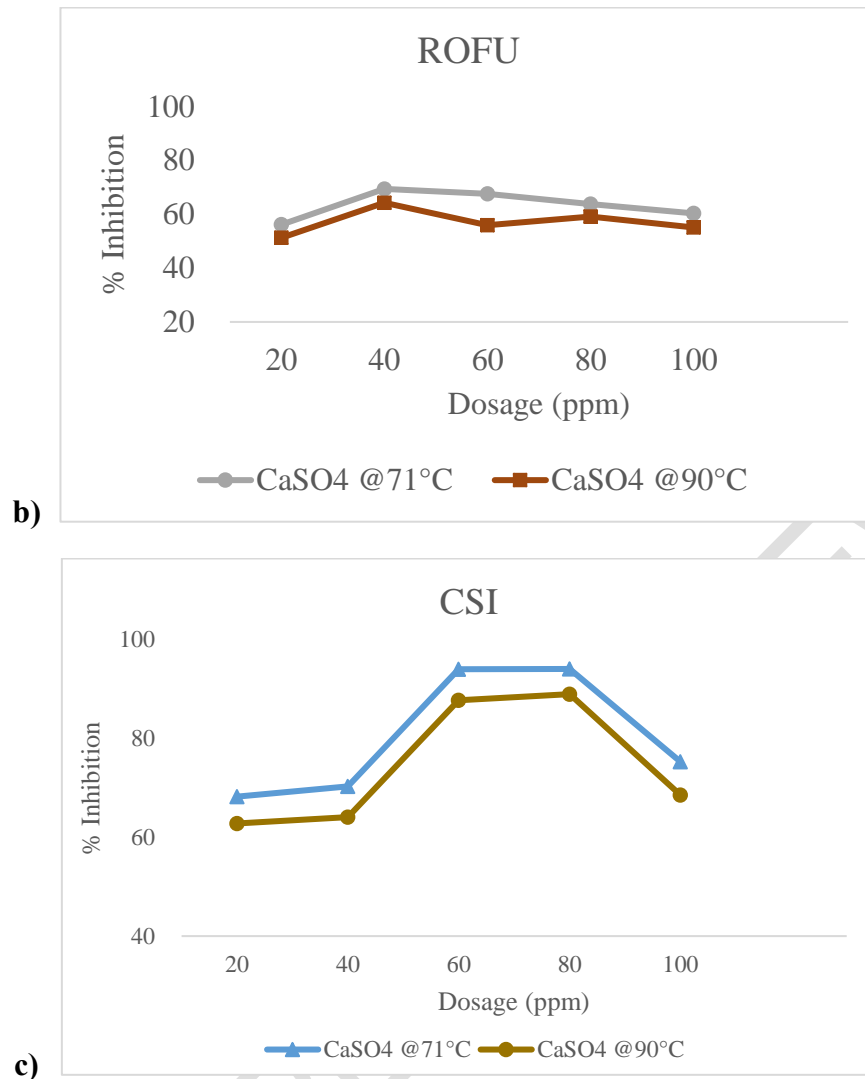
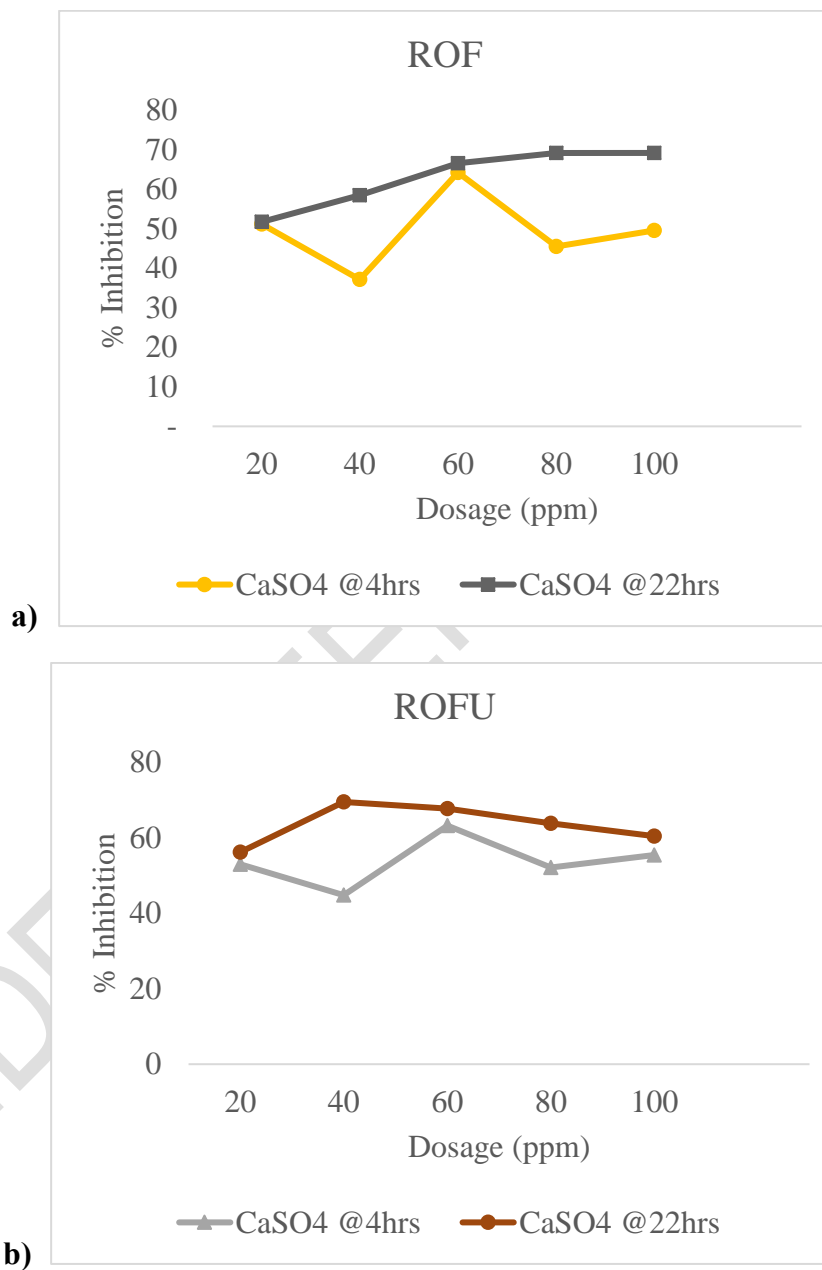


Figure 5 Effect of temperature on the performance of (a) ROF, (b) ROFU, and (c) CSI on CaSO₄ scales at 71°C and 90°C at 22hrs, respectively

3.5.2 Contact time effect

The study examined the impact of contact time on the inhibition of CaSO₄ and CaCO₃ scales at 4 and 22 hours, as shown in Figures 6 and 7, respectively. A similar trend was observed across the inhibitors evaluated for the scale types studied: longer contact time (22 hrs) led to increased inhibition performance, suggesting that the length of inhibitor-scale contact plays an integral role in the continuous inhibition of scales, particularly the CaSO₄ and CaCO₃ scales. The scale inhibition values increased steadily, with optimal values recorded in CaSO₄ scales than in CaCO₃ scales except for CSI, which showed higher inhibition values in CaCO₃ than in CaSO₄ scales. As concentration increased, the effect of contact time on CSI's ability to inhibit performance on the CaSO₄ scale showed some variations over time, as shown in

Figure 6c. However, with the studied scales, the difference in inhibition efficiency (IE) is minimal with extended contact times. Overall, the data suggests that for both scale types and inhibitors evaluated, ROF and ROFU, increasing the contact time between the scale and the inhibitor improves its performance (Victor-Oji *et al.*, 2024).



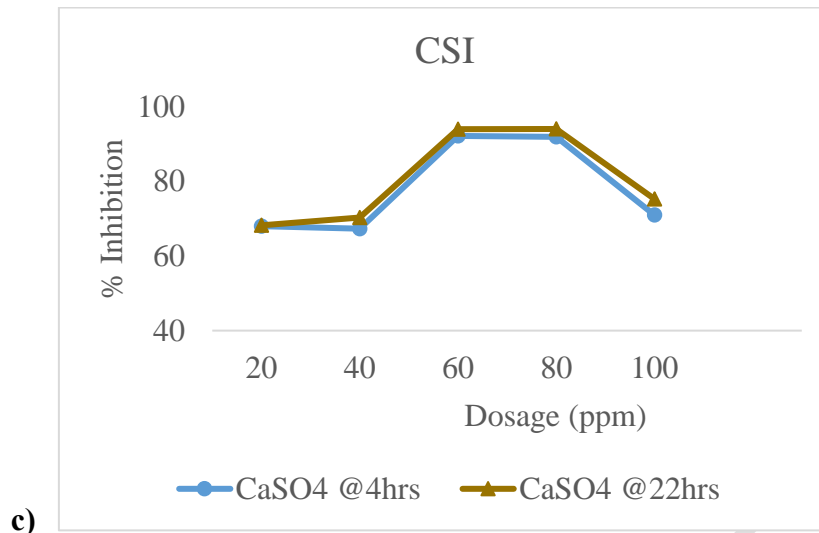
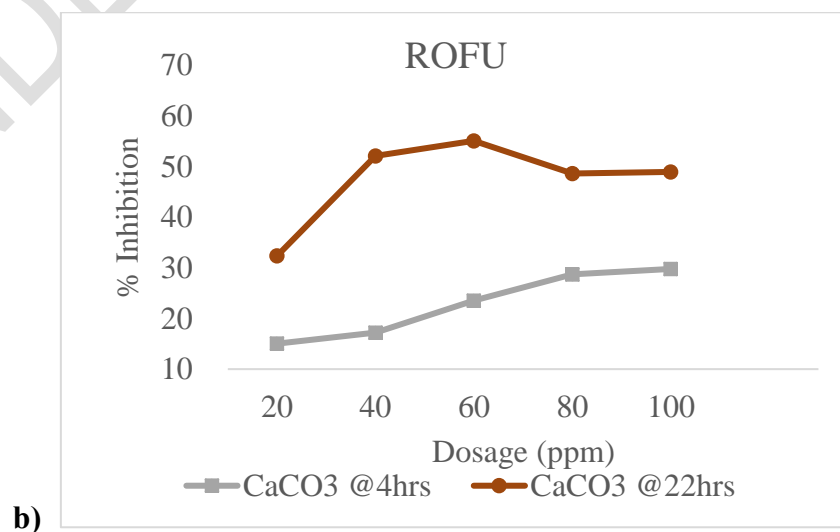
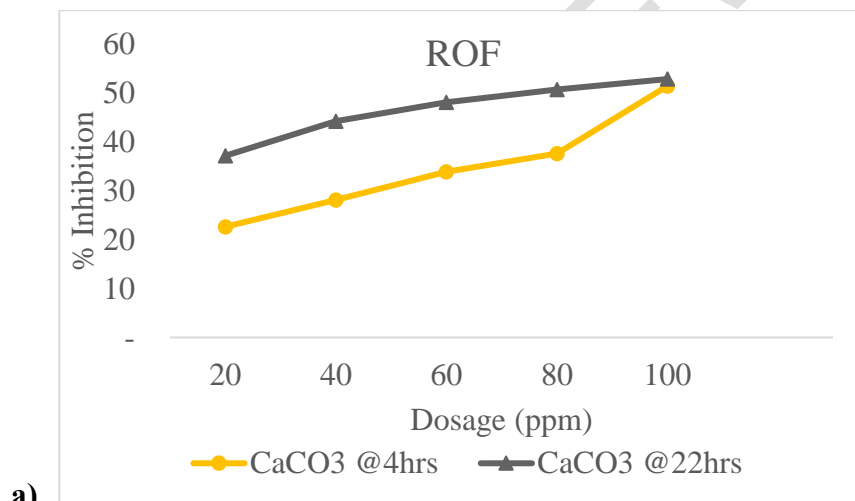


Figure 6 Effect of contact time on the performance of (a) ROF, (b) ROFU, and (c) CSI on CaSO₄ scales at 4 hrs and 22 hrs at 71°C respectively



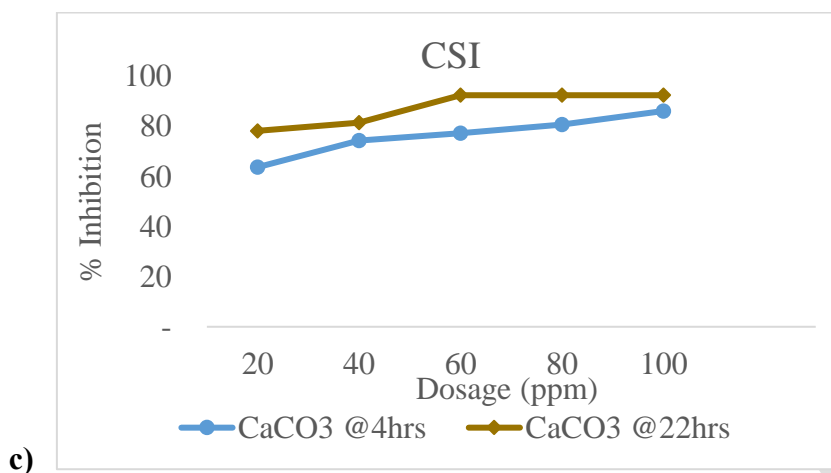


Figure 7 Effect of contact time on the performance of (a) ROF, (b) ROFU, and (c) CSI on CaCO₃ scales at 4 hrs and 22 hrs at 71 °C respectively

3.5.3 Concentration effect

The effect of concentration on scale inhibition using ROF, ROFU, and CSI at different inhibitor dosages (20 ppm to 100 ppm) was evaluated at 71 °C and 90 °C for 22 hours and is shown in Figures 8 and 9 for CaSO₄ and CaCO₃ scales, respectively. Generally, increasing the inhibitor dosage for the studied inhibitors increased the inhibition efficiency (Spinthaki *et al.*, 2021; Una *et al.*, 2021; Victor-Oji *et al.*, 2024). However, once optimum inhibition is achieved, the inhibition rate either remains constant or slowly declines with increasing concentration. A comparative analysis of the formulated SIs with a commercial scale inhibitor, CSI, showed that despite its overall high inhibition rates across the studied scale types, ROF and ROFU show inhibition potential as oilfield green scale inhibitors. ROFU attained optimal inhibition of 69.45% and 55.05 % at 71 °C, 64.30% and 55.85% at 90 °C, and lower concentrations of 40 ppm and 60 ppm, while for ROF, optimal inhibition of 69.15% and 52.86% at 71 °C and 100 ppm, 68.69% and 58.82% at 90 °C, and concentrations of 80 ppm and 100 ppm for CaSO₄ and CaCO₃ scales for 22 hours, respectively.

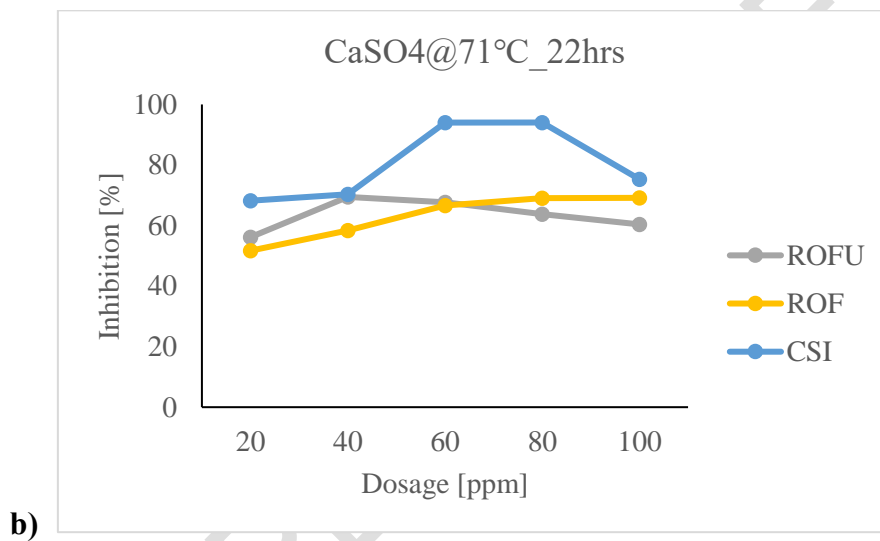
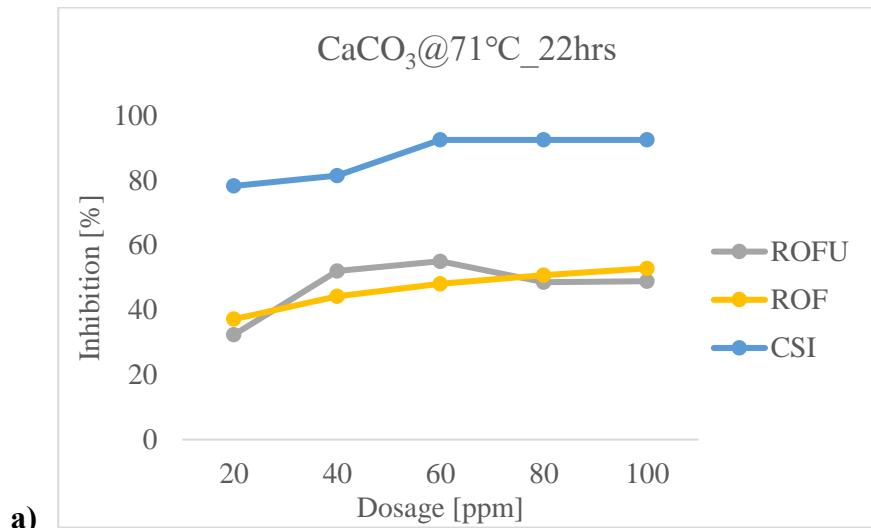
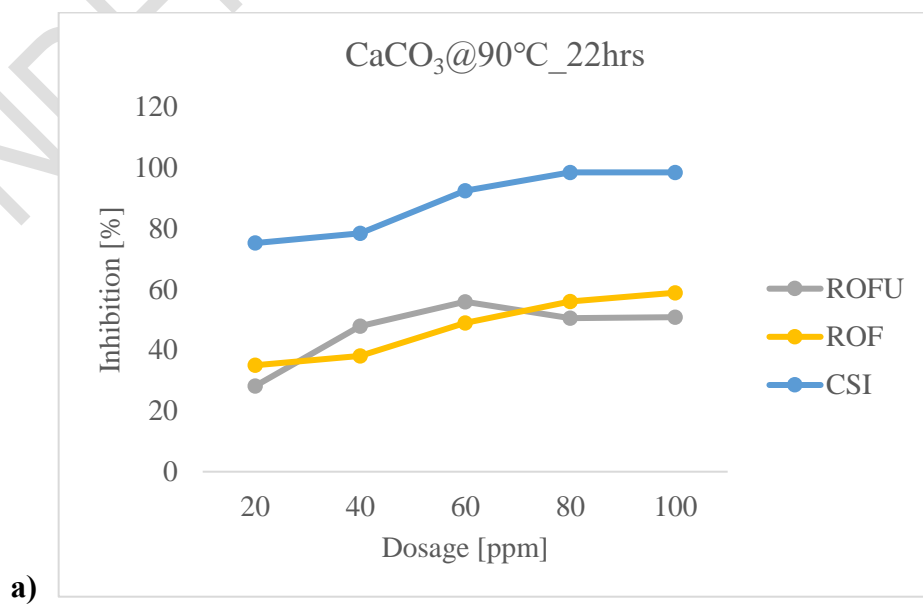


Figure 8 Effect of concentration on the inhibitor performance of (a) CaCO₃ scales and (b) CaSO₄ scales at 22 hrs and 71°C



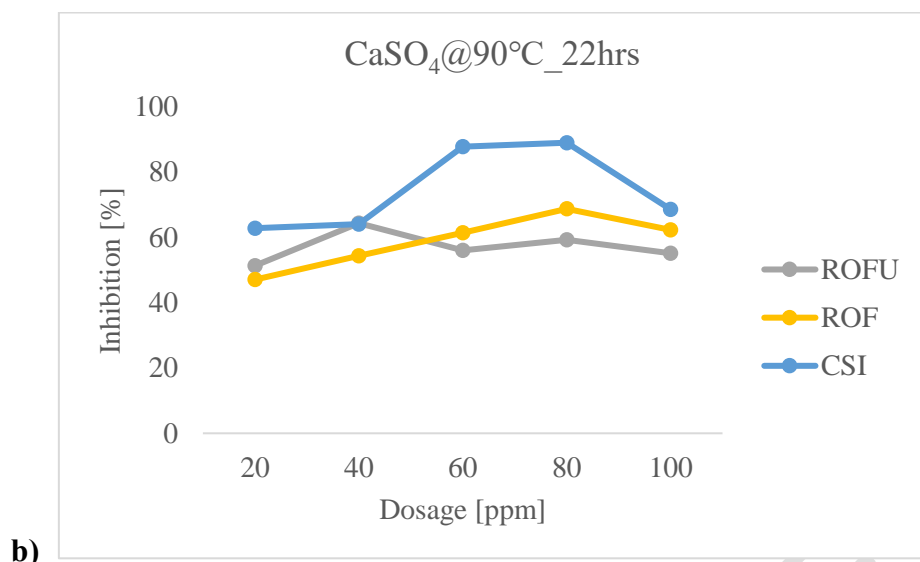


Figure 9 Effect of concentration on the inhibitor performance of (a) CaCO₃ scales and (b) CaSO₄ scales at 22 hrs and 90°C

3.5.4 Effect of Chemical Structure

From the inhibition data of the formulated scale inhibitors, we can infer that ROF effectively inhibited CaCO₃ scales. At the same time, ROFU showed high inhibition values for CaSO₄ scales, with longer contact times and/or increased temperature and concentration dosage. This may be attributed to the increased number of hydroxyl groups introduced into the ROSE moiety during the chemical modification of ROF with urea via the introduction of amide groups (Una *et al.*, 2021).

4. Conclusion(s)

The scale-inhibiting potential of a potential green oilfield scale inhibitor derivatized via chemical modification of red onion skin extract (ROSE) with furfuraldehyde (ROF) and urea (ROFU) on Calcium sulfate and carbonate scales has been evaluated. From compatibility and thermal stability studies, the formulated scale inhibitor blends, ROF and ROFU, were found to be thermally stable, highly soluble, and compatible with the simulated brine. Using the NACE standard static bottle test method, the performance evaluations showed that ROF and ROFU are more effective in inhibiting the growth of CaSO₄ scales than CaCO₃ scales, as moderately high IE of 69.15% and 69.45% for CaSO₄ scales, unlike the lower IE values; 51.42% and 29.77% observed for CaCO₃ scales, at the studied concentration range. From the

inhibition data, we can infer that ROF effectively inhibited CaCO_3 scales. At the same time, ROFU showed high inhibition values for CaSO_4 scales, with longer contact times and/or increased temperature and concentration dosage. This may be attributed to the increased number of hydroxyl groups introduced into the ROSE moiety during the chemical modification of ROF with urea via the introduction of amide groups. Unlike other studies, it was observed that temperature increase had a negligible impact on the inhibition rate for all the inhibitors and both scale types. However, their IE increased with the contact time, as the formulated inhibitors both demonstrated continuous scale inhibition action. Contrary to the CaSO_4 scale, which requires lower temperatures, the CaCO_3 scale requires high temperatures and inhibitor concentrations for effective inhibition. Based on literature findings, increasing inhibitor concentration is required for improved efficiency since higher temperatures promote the development of CaSO_4 scales (Dyer & Graham, 2002; Olajire, 2015; Spinthaki *et al.*, 2021; Una *et al.*, 2021; Victor-Oji *et al.*, 2024). In conclusion, the performance of the formulated SIs, though slightly lower across the concentration range studied for CaCO_3 and CaSO_4 scales, compared favorably with CSI in terms of their ability to inhibit and, as a result, reveal they can be considered potentially inexpensive, renewable, eco-friendly, and effective green oilfield scale inhibitor alternatives for CaSO_4 scales. This enables further research into the industry's application of ROSE derivatives as green oil field scale inhibitors; furthermore, the adsorption mechanism of these inhibitors should be investigated.

Availability of data and materials

All the datasets generated during this study are available on reasonable request.

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