The influence of low amounts on in situ-polymerized bisphenol-diamine net in cement slurries prepared in seawater – structural analysis after long-term contact with in situ-generated mud-acid fracturing fluid
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The influence of low amounts on *in situ*-polymerized bisphenol-diamine net in cement slurries prepared in seawater – structural analysis after long-term contact with *in situ*-generated mud-acid fracturing fluid

Influência de pequenas quantidades do bisfenol-diamina polimerizado *in situ* em pastas de cimento preparadas com água do mar-análise estrutural após longo tempo de contato com fluido de fraturamento *mud acid* gerado *in situ*

Danilo Oliveira Santos¹, Ivory Marcos Gomes dos Santos¹, Joenesson Filip Santos Ribeiro ², Eunice Fragoso Silva Vieira², Gwenn Le Saoût ³, Antonio Reinaldo Cestari¹, ²

¹ Department of Materials Science and Engineering, Federal University of Sergipe, CEP: 49100-000, São Cristóvão, Sergipe, Brazil. e-mail: danilo.quimico@yahoo.com.br, ivorysantos@gmail.com, arcestari@gmail.com
² Laboratory of Materials and Calorimetry, Department of Chemistry/CCET, Federal University of Sergipe, CEP: 49100-000, São Cristóvão, Sergipe, Brazil. e-mail: joenesson.joe@hotmail.com, eunice@ufs.br
³ Lab. de Mécanique et Génie Civil (LMGC), IMT Mines Ales, Univ. Montpellier, CNRS, Ales, France. e-mail: gwenn.le-saout@mines-ales.fr

ABSTRACT

Inadequate zonal isolation in oil wells can provoke safety and environmental problems. New cement slurries can avoid such problems. In this work, novel cement slurries were prepared in the presence of seawater with *in situ*-formed epoxy resin–ethylenediamine. The new slurries were tested in relation to their mechanical properties and characterized by FTIR, XRD, TG/DTG, porosimetry by N₂ adsorption-desorption and solid-state ²⁹Si and ²⁷Al NMR. Some kinetic features of long-term contact of the slurries with *in situ*-generated mud-acid were performed to evaluate the performances of the new slurries in routine acidizing procedures in oil well industry. The experimental kinetic data were well fitted to order-variable exponential kinetic model and a factorial design of four variables (2⁴). In these studies, it was possible to evaluate in detail how some important experimental factors act on the quantities of acid that interact with the pastes, as well as to verify how these factors affect the speed with which the mud-acid interacts with the slurries. The results strongly suggest that the interactions occurred at epoxy/ethylenediamine cement slurry/mud-acid interfaces are due to surface reactions with preservation of the chemical structures of the cement slurries, even after long-term contact with mud-acid. The new cement slurry present good features to be used in environmental-friendly procedures in oil well field.

Keywords: Epoxidized bisphenols; cement slurries; characterization of oil well materials; kinetic modeling; Multivariate analysis.

RESUMO

O isolamento zonal inadequado em poços de petróleo pode provocar problemas de segurança e ambientais. Novas pastas de cimento podem evitar esses problemas. Neste trabalho, novas pastas de cimento foram preparadas na presença de água do mar com resina epóxi-etilenodiamina formada *in situ*. As novas pastas foram testadas em relação às suas propriedades mecânicas e caracterizadas por FTIR, DRX, TG / DTG, porosimetria por adsorção-dessorção de N₂ e RMN de ²⁹Si e de ²⁷Al no estado sólido. Algumas características cinéticas do longo tempo de contato das pastas com o *mud acid* gerado *in situ* foram realizadas para avaliar o desem-
penho das novas pastas nos procedimentos rotineiros de acidificação na indústria de poços de petróleo. Os dados cinéticos experimentais foram bem ajustados ao modelo cinético exponencial de ordem variável e ao planejamento fatorial de quatro variáveis \((2^4)\). Nesses estudos, foi possível avaliar detalhadamente como alguns fatores experimentais importantes atuam sobre as quantidades de ácido que interagem com as pastas, bem como verificar como esses fatores afetam a velocidade com a qual o mud acid interage com as pastas. Os resultados sugerem fortemente que as interações ocorridas nas interfaces da pasta de cimento epoxi / etilenodiamina / mud acid são devidas a reações de superfície com preservação das estruturas químicas das pastas de cimento, mesmo após contato prolongado com mud acid. A nova pasta de cimento apresenta boas características para ser usada em procedimentos nos poços de petróleo.

**Palavras-chave**: Bisfenóis epoxidados; pastas de cimento; caracterização de materiais de poços de petróleo; modelagem cinética; Análise multivariada.

1. **INTRODUCTION**

Among other features, cement slurries have been used to block undesirable underground streams of oil, gas, water or the like [1,2]. Mainly for security reasons, oil well cementing has very low tolerance for failures. In this manner, new-engineered oil well cement slurries have been developed to enable adequate safe and long-term performance of oil and gas wells [2,3]. The so-called primary oil well cementing is made to provide a suitable and stable bond in underground formations, as well as to protect the pipe. Nowadays, cement-based materials have been developed to present chemical/physical stabilities and ensure adequate well durability.

The production levels of wells may be affected by the natural characteristics of the reservoir and damages from routine operations at great depths. In this way, acid fracturing has been a technique in oil industry for solving problems of some oil wells with impaired production. This technique consists in using acidic fluids to crack some rocks in specific underground environments, employing inorganic acid-based fluids, typically mud-acid, a mix of HCl and HF aqueous solutions [4]. However, such routine operations have been understood as an important aspect to oil and gas industries due to possible undesirable long-term reactions between the cement slurry in the annulus section and the acidic fluid, leading to the development of wormholes in some fractured interfaces. In many cases, undesirable chemical reactions result in soluble salts, which are leached away by underground water. From practical point of view, time-related rates of the acidic interactions are of utmost importance to choose a given cement slurry composition for long-term used in oil well field [5]. On the other hand, many molecular-level characteristics of hardened cement slurries remain unclear. It is mainly due to the partial destruction of the structure of cement-based materials after long-term contact with onshore/offshore oil well acidic environments [6-8]. In this way, suitable performances of oil well cement slurries have been reached by the use of some additives [6,7].

Some reactive polymeric additives have been observed to improve the structural features of hardened oil well cement bodies. In addition, the integration of polymer/cement is also known to provide lighten slurries, with better tensile strength, adhesions, and durabilities than cement slurries without such additives. Some polymers, such as styrene butadiene copolymers, PVC, and polyacrylcs have been successfully applied in oil well procedures. Particularly, polymeric systems consisting of epoxy resins and diamines as hardeners have shown adequate aspects to be used in cement slurries formulations [9]. In such type of polymerized material, diglycidyl ether of bisphenol A, which is a product of reaction of bisphenol A and epichlorohydrin, is primarily cured by a given diamine producing a crosslinked polymerized resin, where thermal curing of the polymeric system is not a pre-requisite beforehand [10]. In fact, diamine-extended epoxy polymers have been reported to have excellent reactive characteristics in such polymeric system, where the presence of ether groups, polar aliphatic hydroxyls, and some residual unreacted amine groups provides strong interactions between bisphenol-type polymeric nets and cement [11,12]. Yet, another important advantage is that diamine-extended epoxy resins may be used in relatively low amounts in a given cement formulation (typically less than 10 % in relation to cement mass), without loss of efficiency or remarkable changes of the rheological properties of the cement slurry [13]. On the other hand, the quality of the water used to mix oil well cement is a very important parameter and varies depending upon the technical specifications of cementing procedures.

Seawater has been used as the mix water for squeeze operations in offshore cementing procedures [14,15]. The advantages of the use of seawater in oil well cementing are related mainly to (1) ready availability, (2) relative low cost, since transportation can be eliminated, and (3) reduced fresh-water use and storage on offshore platforms. However, it is known that some components of seawater, such as Cl and SO\(_4\)\(^{2-}\), can cause changes in the compositions and properties of cement slurries. In addition, little is known on the characteristics of cement slurries after interaction with mud-acid.
In this work, we report the preparation of cement slurries additivated with epoxy resin and a diamine in the presence of seawater. The structural features of such material are also evaluated before and after long-term interaction with in situ-generated mud-acid in relation to contact time, initial concentration of the acid fluid and temperature. To the best of our knowledge, this is one of the very limited examples of long-term study of polymer-modified cement slurries in the presence of a little studied acidizing medium of mud-acid.

2. MATERIALS AND METHODS

2.1 Materials

Class A oil well cement and silica were both a free gift from Schlumberger - Petroleum Services Ltd (Nossa Senhora do Socorro, Sergipe State, Brazil), both presenting particles diameters of 300-325 mesh. The A/F bisphenol solution was from The Huntsman Co Special Resins, São Paulo, Brazil), which presents branched epoxidized oxirane rings in its chemical structure. Fig. 1 shows the chemical structures of the bisphenol resin and the hardener. Ethylenediamine, concentrated HCl aqueous solution (both from VETEC/Brazil) and tetrafluoroboric acid (Sigma-Aldrich) were all of analytical degree and used without purification. For comparative porpoises, different slurries preparations were made with double-distilled water and a seawater sample, which was collected near an off-shore oil well station in Aracaju, Sergipe State, Brazil.

2.2 Methods

2.2.1 Preparation of the cement slurries

The cements slurries were prepared as described earlier in literature [16,17], targeting hardened slurries with water/cement and silica/cement ratios of about 0.50 and 0.35, respectively, and densities of about 1.75 ± 0.12 g/cm³ [17,18]. Briefly, the main slurry was prepared with 200 g of cement, 70 g of powder silica, 100 mL of seawater, and a mixture of 5:5 (% in relation to cement mass) of A/F bisphenol solution and ethylenediamine. The cement suspension obtained was mixed at 4,000 rpm for 15s (pre-mixing of the wet and dried components) and 35s at 12,000 rpm for an enough mixing of all components of the wet slurry. The demolded specimens were cast into 5.0 cm side cubic molds and stored in a desiccator saturated with water vapor at room temperature for 3h prior to removal from the molds. The cement specimens were then cured directly in seawater for 28 days at 25°C. Slurry with only cement, silica and seawater, as a control material, was prepared and hydrated using the same procedure. Additionally, the same slurries were also prepared and cured using bidistilled water instead of seawater. For simplicity, the cement slurries are hereafter denominated as slurry-standard-SW (cement/silica/seawater), slurry-standard-W (cement/silica/bidestilled water), slurry-MOD-SW (cement/silica/polymeric modifier, seawater) and slurry-MOD-W (cement/silica/polymeric modifier, bidestilled water). Samples were carefully removed from the surface of the cemented specimens and characterized.

2.2.2 Long-term immersion tests in in situ-produced mud-acid

The immersion experiments were made by isothermal procedures from 25 to 55 ± 0.1 °C for approximately 90 hours, using mud-acid aqueous solutions from 0.01 to 0.1 molL⁻¹ in relation to their total acidic content of the solutions. The mud-acid was a mixture of 12% HCl + 3% HF, where the HF was in situ-generated from a reaction of tetrafluoroboric acid in water [19]. In a typical experiment at a given temperature, 50 mL of mud-acid were added to 100 mg of determined slurry in a 150 mL polyethylene flask and continuously stirred. Samples were taken at predetermined times and the total acidic content of the solution was determined by a
pH-meter or acid–base titrations using 0.01 mol L\(^{-1}\) NaOH solutions. All experiments were carried out in triplicate runs. The amounts of total acidic species interacted with the slurries in relation to contact time (\(Q_t\)) were calculated using the expression [20]:

\[
Q_t = \frac{(c_i-c_f)V}{m}
\]

(1)

where \(Q_t\) is the interacted amount of acidic species per gram of slurry in mol g\(^{-1}\), \(c_i\) is the initial concentration of acidic species in solution, in mol L\(^{-1}\), \(c_f\) is the concentration of acidic species after a given contact time, in mol L\(^{-1}\), \(V\) is the volume of the solution in L, and \(m\) is the mass of slurry in g.

A factorial design study was also conducted to determine principal and interaction effects of selected experimental factors of mud-acid interaction onto the slurries. Such types of effects were calculated using Eq. 2:

\[
Effect = \bar{R}_{+,-} - \bar{R}_{-,+}
\]

(2)

where \(\bar{R}_{+,-}\) and \(\bar{R}_{-,+}\) are average values of \(Q_t\) for the high (+) and low (−) levels of each factor. For principal effects, the above averages simply refer to the results at high (+) and low (−) levels of a given factor, which was calculated regarding the levels of the other factors. For binary interactions, is the average of results for the factors at their high and low levels, whereas is the average of the results for which one of the factors involved is at the high level (+) and the other is at the low level (−). High-order interactions are calculated using Eq (2), applying signs obtained by multiplying those for the factors involved for high and for low levels. Since duplicate runs were performed for each individual factorial design experiment, standard errors (E) of the effects were calculated by Eq. (3):

\[
E = \left\{\sum(d_i)^2/8N\right\}^{1/2}
\]

(3)

where \(d_i\) represents the difference of each duplicate value and \(N\) is the number of the experiments performed.

### 2.2.3 Characterization of the samples

Infrared spectral spectra were obtained on a Varian FTIR spectrophotometer, model 640 using KBr pressed-disks technique at room temperature was used in the spectral range from 4000 to 900 cm\(^{-1}\). Thermal analysis curves (TG/DTG) were obtained with about 10 mg of a given slurry under nitrogen atmosphere at 10 °C min\(^{-1}\) from 25 to 900°C, in a SDT 2960 thermoanalyzer from TA Instruments. The X-ray diffraction (XRD) diffractograms of the samples were obtained in a Shimadzu LABX 6000 diffractometer (CuK\(\alpha\) radiation; \(\lambda = 1.5400\) Å, 2\(\theta = 5\) - 60°), with accumulation rate of 0.02° min\(^{-1}\). The porosity characteristics of the cement slurries were determined by N\(_2\) adsorption-desorption experiments performed at 77K on a NOVA 1200 analyzer. The specific surface area (SBET) was determined from the Brunauer-Emmet-Teller (BET) equation. Pore size distributions were estimated using the Barrett-Joyner-Halenda method.

The \(^{29}\)Si and \(^{27}\)Al MAS NMR spectra were obtained using a Bruker Avance 400 NMR spectrometer using ZrO\(_2\) rotors, as described earlier [17]. The \(^{29}\)Si chemical shifts were referenced to tetramethylsilane (TMS) at 0.0 ppm and the \(^{29}\)Si isotropic chemical shifts were resolved and measured to ± 0.1 ppm. The field strength was of 9.4 T at 79.49 MHz, with 5 kHz spinning rates on a 7.5 mm CP/MAS probe, with single-pulse experiments being carried out by applying 900 pulses with 1H decoupling and recycle delays of 10 s. The \(^{27}\)Al MAS NMR spectra were recorded on a Bruker Avance III 600 NMR spectrometer. The \(^{27}\)Al chemical shifts were referenced relative to a 1.0 mol L\(^{-1}\) AlCl\(_3\)-6H\(_2\)O solution at 0.0 ppm. The field strength was 14.1 T at 156.33 MHz, with 20 kHz spinning rates on a 3.2 mm CP/MAS probe using also ZrO\(_2\) rotors. Single-pulse experiments were carried out by applying π/12 pulses and recycle delays of 1s.

Mechanical tests of the specimens were carried out on an Instron Universal Testing Machine, model 3385H. The compressive strength test used a 250 kN load cell with a loading speed of 0.5 mm.min\(^{-1}\). The moduli of elasticity of the samples were obtained from slope of the linear region of the load-displacement curve.
3. RESULTS

3.1 Characterization of the cement samples

Characterization has been an important tool for understanding the relationship chemical composition-molecular structure associated of hydrated cement slurries. Furthermore, characterization is intended in this work to provide additional molecular-level information after the cement slurries were in long-term contact with chemically aggressive acidic media. Cement slurries are indeed a multiphase and complex class of materials.

Typically, there are four main mineral phases in oil-well cements [1]: tricalcium silicate (C₃S, Ca₃(SiO₅)₂), dicalcium silicate (C₂S, Ca₂(SiO₄)₂), tricalcium aluminate (C₃A, Ca₃(Al₂O₆)), and calcium aluminoferrite (C₄AF) of different compositions. Upon hydration, the calcium silicate hydrate (C–S–H) is formed, being a relatively poorly crystalline material consisting firstly of nonstoichiometric dimeric units, which present slow polymerization to give higher linear pentamers and octamers. A number of methods has been used to characterize amorphous and crystalline components of cements.

Figs. 2 and 3 show the FTIR spectra of the slurries, before and after mud-acid interaction. However, analysis of some bands in the mid-IR spectra (2800–4000 cm⁻¹) is indeed difficult due to the presence of broad bands and bands overlaps. Anyway, this technique enables to find some features of physically adsorbed water, silicate polymerization, possible presence of Ca(OH)₂ (CH, portlandite) and TO₄-type condensed tetrahedral in cement chains (T = Si or Al in Si-O-Si or Al-O-Si bonds), as well as possible presence of calcium carbonate polymorphs [21].

For the standard slurries (Fig. 2, a and c)), the small narrow bands at 3,640 and 3,642 cm⁻¹ are due to CH (Ca-OH vibrations). The broad bands centered at 3446 and 1690 cm⁻¹ are due to O-H groups in H₂O or hydroxyls groups of hydrated products, which present several hydrogen bond strengths [22]. The band centered at around 1081 cm⁻¹ is assigned to Si-O stretching vibrations of the SiO₄ tetrahedral units, from residual unreacted silica or C-S-H mixes [23]. Condensation reactions between Al-OH and Si-OH groups, forming aluminosilicate bonds, can also occur. The Si-O stretching bands, however, usually overlap ettringite (Ca₆Al₂(SO₄)₃(OH)₁₂.26H₂O) bands at around 1000 cm⁻¹ and 1120 cm⁻¹. The band at 453 cm⁻¹ is associated with bending vibrations of O-Si-O groups, which is influenced by O-Si-O angles or influence of neighboring sites. Bands of calcium carbonate can be found at about 1420 and 850 cm⁻¹. Typically, the presence of crystal structures of calcium carbonate polymorphs (vaterite, aragonite and calcite) can be made using FTIR. In this manner, the small bands centered at 873 and 1427 cm⁻¹ suggest the presence of aragonite [24]. The mid-IR bands changed in frequency, which are related with silicate reactions after mud-acid interaction [25,26]. The bands at 3640 and 1086 cm⁻¹ are not clearly observed after mud-acid interaction suggesting, respectively, partial reactions of leaching of CH and depolymerization with rearrange of silicates chains, leading to formation of new aluminosilicates [27].

Figure 2: FTIR spectra of the standard slurries: a) Slurry-Standard-W; b) Slurry-Standard-W after mud-acid interaction; c) Slurry-Standard-SW and d) Slurry-Standard-SW after mud-acid interaction.
Regardless of the characterization technique used, the identification of polymers within cement-based matrixes has been a major problem, due to the low amounts of such compounds in cement samples. Looking at the FTIR spectra of the chemically modified cement materials (Fig.3) the formation of epoxy/ethylenediamine polymeric net seems to cause reorganizations of hydrated cement microstructure. The presence of very small broad bands at 2925 and 1795 cm$^{-1}$ are due to both alkyl (-CH$_3$, -CH$_2$) and ester (-COOR) groups of the epoxy resin, respectively. At 3643 cm$^{-1}$ only a shoulder appears, which may correspond to interactions (probably hydrogen bondings) between hydrated cement phases and the epoxy-ethylenediamine polymeric net [28, 29]. This may be occurred due to the inhibitive effect of the epoxy-ethylenediamine resin on the formation of CH during the hydration of the slurry-MOD. The FTIR spectrum of slurry-MOD after exposure to mud-acid is more or less unchanged in relation to the pristine slurry-MOD. It seems to be a first evidence of the effect of the Epoxy-Ethylenediamine resin (Epoxy-EDA) against the leaching out some chemical components of the polymer-modified cement slurry in the presence of mud-acid.

The XRD plots of the slurries are presented in Figs. 4 and 5. XRD powder diffractions of hydrated cement materials are related to identification of phase composition, possible determination of polymorphic modifications and analysis of crystallinity of individual phases. Some products, however, do not present clear diffraction peaks. In addition, some XRD peaks from different components may coincide or overlap.
The diffractograms of the standard slurries prepared in seawater and distilled water (slurry-standard-SW and slurry-standard-W) show similar behavior one another, in which the presence of the most common hydrated compounds in the cement slurries is verified. For these slurries, the principal peaks are (in 2θ, degree): 9°, ettringite; 18°, CH; 21°, silica; 26°, C-S-H and 29°, calcite [30].

The main products of cement hydration are ettringite, CH and C-S-H. The hydration of the calcium silicates produces a family of structurally similar, essentially amorphous or little crystalline silicates. CH is...
the second most abundant hydration product, forming crystalline aggregates [31,32]. Ettringite is formed by hydration of alita (C₃S) in the presence of gypsum (hydrated calcium sulfate). The formation of calcite (CaCO₃) is attributed to the partial carbonation of CH and presence of carbon dioxide dissolved in water [33]. The peak at 28° refers to a mixture of quartz and calcite in the cement slurries [34]. The presence of C₃S (32°) in the diffractograms suggests that hydration was not complete.

Some aspects of microstructure of the standard slurries have changed after interaction with mud-acid. From analysis of the standard slurries XRD diffractograms, the crystalline aspect of these cement slurries decreased markedly (Fig. 4), where there are no peaks of the mains hydrated phases (CH, C-S-H and ettringite). Some features of chemical reactions of hydrated cement in the presence of acidic solutions have already been studied in detail. Under acid pH, the constituents (CH, C-S-H, calcium aluminate hydrates) dissolve at various velocities over contact time and new salts precipitate. The first and faster dissolving component is CH along the neutralization reactions. In these chemical reactions, there is the formation of water-soluble calcium salts. The strength of the attack is considered mainly dependent on the solubility of the precipitated salts. After leaching the CH to the solution, the matrix C-S-H is attacked. Under appropriate conditions, the chemical attack may result in the complete decomposition of the hydration products [35]. Thus, the degradation of the constituents of these slurries with possible restriction to the added silica is evidenced.

Figure 5 shows the diffractograms of modified slurries before and after the interaction with mud-acid. The diffractograms of the slurry-MOD-SW and slurry-MOD-W did not exhibit additional peaks when compared to the standard slurries. The results of X-ray diffraction analysis show that there is no significant difference between the hydration products of standard slurry and modified slurry with Epoxy-EDA as predicted in the literature [36]. Slurry-MOD presented the following main peaks: 9° ettringite, 18° CH, 21° silica, 26° C-S-H and 29° Calcite. The diffractograms of the slurries, after interaction with mud-acid are similar to those materials before interaction with mud-acid. For these materials, there is a decrease of intensity in the peaks of CH, confirming data from the literature that this phase undergoes preferential decomposition [37]. The peak at 28° for the quartz-calcite mixture is highlighted in the diffractograms of these modified cement slurries, and still found after mud-acid attack. Therefore, it is verified that the presence of Epoxypolymer protected quartz from chemical degradation. The microstructures of the polymer-modified slurries remain relatively unchanged, even after the interaction with mud-acid.

The results from the TG/DTG tests are shown in Figs. 6 and 7. Thermal analysis can be used to quantitatively determine mass loss of liquids (typically H₂O), gases, as well as to indicate the presence of some cement phases, which do not evidenced using XRD technique, because this technique only shows slurries crystalline phases. However, the identification of mineral components and their quantification is a difficult task due to temperature-related overlapping processes [38].

The TG/DTG plots profiles help to evaluate the typical reactions occurring in the cement-based slurry when subjected to a progressive temperature increase. Figure 6 shows the TG/DTG plots of standard slurries. Standard slurries (Slurry-Standard-SW and Slurry-Standard-W) have shown similar TG/DTG curves one another. Typically, there are three main mass losses regarding to the release of evaporation of physically adsorbed water on the surfaces and pores of slurries, as well as to the decomposition of hydrated calcium silicate and ettringite (60-200°C), the decomposition of CH (410-500°C) and the calcination of amorphous materials and calcite (CaCO₃), according to equation (4) [24]:

$$\text{CaCO}_3 (s) \rightarrow \text{CaO (s)} + \text{CO}_2 (g) \quad (4)$$
Figure 6: TG and DTG plots of the standard slurries before (above) and after (below) mud-acid interaction.

Figure 7 shows the TD/DTG curves of modified slurries with the Epoxy-EDA polymer (Slurry-MOD-SW e Slurry-MOD-W). The TG/DTG curves of these materials present five main mass losses, with some mass loss overlays. The mass loss located below 100°C is attributed to the removal of physically adsorbed water in the cement slurry and the event in 160°C refer to the decomposition of C-S-H and ettringite. The event between 280-400°C refers to the decomposition of the Epoxy-EDA polymer. The mass loss located between 410 and 500°C is attributed to the CH decomposition and two overlapping events between 710-800°C are characteristic of the calcination of amorphous materials and calcite [39,40].
Figure 7: TG and DTG plots of the modified slurries before (above) and after (below) mud-acid interaction.

The TG/DTG plots of the standard slurries after mud-acid interactions present distinct profiles of data before contact. The percentage of mass loss of these slurries was approximately 2%. In the TG/DTG curve of the Slurry-Standard-SW after acid contact show mass loss around 330°C attributed to the presence of Fe₂O₃ produced from tetracalcium ferro-aluminate during the cement hydration reaction [41]. However, the TG/DTG curves of Slurry-MOD after exposure to mud-acid solution is more or less the same in relation to the modified slurry before mud-acid interaction.

The CH content were estimated from the weight loss measured from each TG plots between the initial and final temperature of a given mass loss process corresponding TG peak by considering the following decomposition reaction (410-500°C), according to Eq. 5 [32]:

\[
\text{Ca(OH)}_2 (s) \rightarrow \text{CaO (s)} + \text{H}_2\text{O (g)}
\]  

(5)
From TG/DTG curves, the portlandite content was determined using Eq. (6) [32]. Table 1 shows the values of the percentage of CH in each cement slurry.

\[
CH (\%) = \frac{\Delta CH(\%) \cdot MM_{CH}}{MM_W}
\]  

where \(CH (\%)\) is content portlandite, \(\Delta CH(\%)\) is the weight loss during the decomposition of CH, \(MM_{CH}\) is the molar weight of CH and \(MM_W\) is the molar mass of water.

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>CH (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slurry-Standard-SW</td>
<td>11.41</td>
</tr>
<tr>
<td>Slurry-Standard-SW-Mud-Acid</td>
<td>Not found</td>
</tr>
<tr>
<td>Slurry-Standard-W</td>
<td>14.58</td>
</tr>
<tr>
<td>Slurry-Standard-W-Mud-Acid</td>
<td>Not found</td>
</tr>
<tr>
<td>Slurry-MOD-SW</td>
<td>6.97</td>
</tr>
<tr>
<td>Slurry-MOD-SW-Mud-Acid</td>
<td>5.15</td>
</tr>
<tr>
<td>Slurry-MOD-W</td>
<td>6.97</td>
</tr>
<tr>
<td>Slurry-MOD-W-Mud-Acid</td>
<td>4.64</td>
</tr>
</tbody>
</table>

Standard slurries have higher amounts of CH than the modified slurries with Epoxy-EDA polymer. A possible explanation may be the interaction of the polymers at the CH nucleation sites, which prevents the growth of their crystals [32]. The presence of polymers can significantly influence the morphology of CH in cement matrices, depending on the type of polymer, the proportion of polymer in the cement or a combination of all these factors [42]. After the mud-acid attack, the amount of CH was calculated only for the modified slurries, since for the standard slurry no thermal events of mass losses attributed to the presence of CH were identified. For modified slurries, the percentage of CH was lower after the acid attack suggesting partial destruction of the CH structures in the slurries containing the Epoxy-EDA polymer.

It has been shown that the presence of Epoxy-EDA can significantly influence the properties of cement slurries. It was evidenced that modified slurries presented superior behavior in relation to the standard slurries against to acid attack, suggesting that the polymer directly influenced the resistance of the slurries. These all conclusions from TG/DTG curves seem to be very consistent with results from FTIR and XRD characterization.

![Figure 8: $^{29}$Si MAS NMR spectra of the modified slurries: a) Slurry-MOD-W; b) Slurry-MOD-W after mud-acid interaction; c) Slurry-MOD-SW and d) Slurry-MOD-SW after mud-acid interaction.](image-url)
In general, $^{29}$Si chemical shifts in silicates are located from -60 to -120 ppm. In this work, the $^{29}$Si chemical shifts were analyzed using the $Q_n(m\text{Al})$ classification, where one given Si tetrahedron is connected to $n$ Si tetrahedra ($n = 0 – 4$); $m$ is the number of neighboring AlO$_4$ tetrahedra [43]. The $^{29}$Si NMR spectra of modified slurries, before and after contact with mud-acid are shown in Figure 8. The $^{29}$Si NMR spectra of Slurry-MOD-SW and Slurry-MOD-W are similar. The spectra show near -70 ppm resonances from the $Q_0$ units of the anhydrous cement (unreacted cement). The broad peaks with maxima at -78 and 84 ppm are due to $Q_1$ and $Q_2$ units refer to polymeric end-chain tetrahedral and middle nonbridging tetrahedral units in polymer chain of C-S-H, respectively. The small broad peak of $Q_4$ type site centered at -106 ppm is related to crystalline quartz [43,44].

After contact with mud-acid, a small and broad $Q_1$ peak is observed near -98 ppm, which is referred to formation of highly polymerized amorphous alumino silicate phases [45]. The effect of mud-acid is more destructive for the C-S-H, mainly silica gel is present in both samples. C-S-H seems to be more resistant in the Slurry-MOD-SW. Interestingly, the $Q_4$ peak remains in the $^{29}$Si NMR spectra, corroborating with the analyzes of XRD that indicate the presence of quartz after the contact with mud-acid solution [43].

$^{27}$Al NMR is used to differentiate Al in different coordination states in the structural characterization of aluminate phases in cement slurries [46]. $^{27}$Al NMR data recorded are shown in Fig. 9. The $^{27}$Al NMR spectra of the modified slurries are quite similar. These spectra shown Al in fourfold coordination due to Al substituting Si in the C-S-H. In the aluminum sixfold range, three main peaks are observed: ettringite at 13 ppm, Calcium Aluminate Monosulfate (AFm), around 9 ppm, and a small shoulder at around 3.5 ppm, which has been related to an amorphous aluminate phase, from precipitation reactions at the surface of a C-S-H or found as a separate phase (TAH) [46,47]. After interaction with mud-acid, Slurry-MOD-W show resonances from tetrahedrally coordinated Al (40-70 ppm), present amorphous alumino silicate phases and in range -10 to 20 ppm can be attributed to octahedrally coordinated Al and third aluminate hydrate [47]. $^{27}$Al NMR spectra of the Slurry-MOD-SW after contact with mud-acid is similar to Slurry-MOD-W with additional chemical shift value attributed to AFm at 9 ppm.

![Figure 9](image-url)  
**Figure 9:** $^{27}$Al MAS NMR spectra of the modified slurries: a) Slurry-MOD-W; b) Slurry-MOD-W after mud-acid interaction; c) Slurry-MOD-SW and d) Slurry-MOD-SW after mud-acid interaction.
For standard slurries, the process of acidic attack resulted in the total decomposition of the hydration products and silica as shown in the previous characterizations (details not shown). Similar results were also observed earlier [17].

**Figure 10:** Adsorption-desorption isotherms of nitrogen at 77K of standard slurries (above) and modified slurries (below).
The features related to the porosity of the cement slurries are considered important to evaluate the durability of cement slurries. The porous structure has a direct effect on the chemical and mechanical resistance of cement slurries [48].

The porosity analysis of the cement slurries was performed, before and after an interaction with the mud-acid. The adsorption/desorption isotherms of the slurries before and after interaction with mud-acid solution are shown in Fig. 10. The cement slurries exhibited a Type II adsorption isotherms, according to IUPAC classification [48].

The isotherms of the standard slurries, after the acid attack, presented different profile in relation to the initial isotherm. After the interaction of these slurries with mud-acid solution, the isotherms showed hysteresis, suggesting that the nitrogen adsorption and its desorption process occurred in different ways, due to changes in the geometry and pore size of the slurries, as shown in Table 2. For modified slurries, the type II isotherm profile was maintained after contact with mud-acid solution, and the geometry and pore size remaining unchanged in these materials.

Table 2: Textural proprieties of the cement slurries.

<table>
<thead>
<tr>
<th>MATERIAL</th>
<th>SURFACE AREA (m².g⁻¹)</th>
<th>PORE VOLUME (cm³.g⁻¹)</th>
<th>PORE SIZE (Å)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slurry-Standard-W</td>
<td>91.5</td>
<td>0.081</td>
<td>18.0</td>
</tr>
<tr>
<td>Slurry-Standard-W-Mud-acid</td>
<td>147.7</td>
<td>0.156</td>
<td>22.6</td>
</tr>
<tr>
<td>Slurry-MOD-W</td>
<td>11.8</td>
<td>0.024</td>
<td>18.2</td>
</tr>
<tr>
<td>Slurry-MOD-W-Mud-acid</td>
<td>35.8</td>
<td>0.076</td>
<td>18.1</td>
</tr>
<tr>
<td>Slurry-Standard-SW</td>
<td>74.6</td>
<td>0.072</td>
<td>18.0</td>
</tr>
<tr>
<td>Slurry-Standard-SW-Mud-acid</td>
<td>101.2</td>
<td>0.141</td>
<td>22.3</td>
</tr>
<tr>
<td>Slurry-MOD-SW</td>
<td>17.6</td>
<td>0.030</td>
<td>18.2</td>
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<tr>
<td>Slurry-MOD-SW-Mud-acid</td>
<td>46.5</td>
<td>0.073</td>
<td>18.3</td>
</tr>
</tbody>
</table>

Modified slurries have pore diameter values identical to those of standard slurries. However, the Slurry-MOD showed smaller surface area and pore volume. These data suggest that crosslinked Epoxy-EDA polymeric net may reduce the porosity of the slurry and may have influence how the slurry interacts in the presence of acidic aggressive media. Epoxy-EDA polymer can occupy voids in the microstructure of the cement slurry, reducing permeability, ion diffusion and also improving mechanical resistance (Table 2) [48,49].

The mud-acid interactions changed some textural properties of standard slurries. However, for Slurry-MOD, these properties have undergone small observable variations [48,49]. The surface area, pore volume and pore diameter of the standard slurry increased in size after interaction with mud-acid. For the modified slurries, these changes were less intense, since their pore diameters remained almost unchanged, suggesting that the degradation of these slurries was lower than for the control slurries.

Figure 11: Effect of the interaction of mud-acid on compressive strength of the cement slurries.
Compressive strength results of the cement slurries are shown in Figure 11. Before the interaction of the cement slurries with mud-acid, the compressive strength has similar values with higher values for the modified pastas. However, after the acid attack, the standard slurries presented a greater decrease than the modified slurries, indicating that the Epoxy-EDA slurries have superior resistance to compression. These results are in good agreement with those obtained by Aggarwal et al (2007) [50]. Those authors showed that the addition of polymers to cement slurries increase compressive strengths and decrease ion penetration, reducing the deterioration of cement slurries. It can conclude that the compressive strength decreases after contact with the mud-acid. However, for standard slurries, these values decrease in greater proportion (approximately 50%), suggesting a better mechanical resistance for modified slurries.

According to Chen et al (2013) [51], the compressive strength is directly related to the porosity of the material. Cement slurries with higher porosity tend to have lower mechanical strength, as presented in this work. Modified slurries exhibit lower porosity, resulting in a better performance in relation to their mechanical properties, before and after the acid attack.

The modulus of elasticity of each cement slurries is shown in Figure 12. It is obvious that the modulus of elasticity decreases after mud-acid interaction. However, the addition of Epoxy-EDA polymer improves the modulus of elasticity for cement slurry. The results of the modulus of elasticity are in agreement with literature. Reis (2010) [52] reported that the decrease of the modulus of elasticity of cement slurries after contact with acidic solutions. According that author, the modulus of elasticity also represents a degradation pattern of the material, in which high values indicate less degradation of the cement slurry. In this way, the Figure 12 indicates that the modified slurries presented less chemical degradation against the acid attack than the standard slurries. Therefore, the in situ-formed branched Epoxy-EDA polymeric net seems to have an efficient potential to mitigate the effects of deleterious processes occurring by the use of API fracturing fluids (mud-acid), during routine acidizing procedures in oil and gas industry.

3.2 Kinetic of mud-acid interaction onto the slurries

During an acidic attack process, the amount of acid consumed is proportional to the amount of cement in the materials attacked, or more precisely to the content of the hydration products, such as CH. Typically, the rate of acid attack decreases with increasing cement content. Therefore, the neutralization capacity of the cement matrix is evaluated [53]. Figure 13 shows the amounts of acid that interacted with the cement slurries at different temperatures. The amount of acid that reacts increases with the contact time and remains constant after a given contact time (approximately 24 hours), when the interaction equilibrium occurs. As shown, the amount of mud-acid interacting with the cement slurries increases with increasing temperature from 25 to 55 °C. Typically, temperature increase enhances acid etching to cement [35].
Many attempts have been made to formulate general mathematical expressions, which would be able to adequately describe the kinetics of interaction at solid/solution interfaces. For efficient analysis of the acid/cement interactions, it is necessary to use suitable mathematical models to calculate the kinetic parameters of the interactions systems. In this work, a kinetic order-variable exponential function was used, as shown in eq. 7 [54]:

\[ Q_t = Q_e \cdot (1 - \exp^{-k\cdot t})^n \]  

(7)

where \( k \) is the kinetic rate (h\(^{-1}\)) and \( n \) is a constant related to the interaction mechanisms.

Figure 13: Experimental results (points) and calculated using the kinetic model of variable order (lines) of the interaction of mud-acid with cement slurries. Initial concentration of \( H^+ = 0.10 \) mol.L\(^{-1}\).

Figure 13 shows the confrontation of the experimental points with the values obtained from the mathematical model, represented by the lines. The comparison of these data shows that this model was well suited to describe the kinetics of interaction between mud-acid solution and cement slurry. The calculated kinetic parameters are shown in Table 3.

In order to evaluate the fit of the kinetic model, the linear least-squares method (\( R^2 \)) was applied. However, \( R^2 \) is sensitive to extreme experimental data points of the correlations plots, resulting in misleading indication of the quality of a given fit. In this work, the fittings of the kinetic model were done using chi-square (\( \chi^2 \)) analysis, according to eq. 8 [32]:

\[ \chi^2 = \sum \frac{(Q_e - Q_m)^2}{Q_m} \]  

(8)

where \( Q_e \) and \( Q_m \) are the acidic amounts at a given time \( t \), which were calculated using experimental data and the kinetic model, respectively. The chi-square statistic test is the sum of the squares of the differences between the experimental data and theoretically predicted data from models. If the calculated data are consistent with the experimental ones, chi-squares will have a small value (generally less than 0.01) [20]. The analysis of the \( \chi^2 \) values suggests that the interaction of the mud-acid with slurries can be described by order-variable exponential model (\( \chi^2 \) values are lower than 4.31x10\(^{-3}\)).
Table 3: Kinetic parameters of the interaction of cement slurries with mud-acid.

<table>
<thead>
<tr>
<th>SLURRY-STD-W</th>
<th>Temperature (°C)</th>
<th>Q.10^{-3} (mol.g^{-1})</th>
<th>SD.10^{-6}</th>
<th>k (h^{-1})</th>
<th>SD</th>
<th>n</th>
<th>SD</th>
<th>R²</th>
<th>X².10^{3}</th>
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<td>0.809</td>
<td>0.020</td>
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<th>k (h^{-1})</th>
<th>SD</th>
<th>n</th>
<th>SD</th>
<th>R²</th>
<th>X².10^{3}</th>
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<tr>
<th>SLURRY-STD-SW</th>
<th>Temperature (°C)</th>
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<th>SD.10^{-6}</th>
<th>k (h^{-1})</th>
<th>SD</th>
<th>n</th>
<th>SD</th>
<th>R²</th>
<th>X².10^{3}</th>
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<td>0.019</td>
<td>0.996</td>
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<td>0.706</td>
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<td>0.732</td>
<td>0.008</td>
<td>0.999</td>
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<table>
<thead>
<tr>
<th>SLURRY-MOD-SW</th>
<th>Temperature (°C)</th>
<th>Q.10^{-3} (mol.g^{-1})</th>
<th>SD.10^{-6}</th>
<th>k (h^{-1})</th>
<th>SD</th>
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<th>R²</th>
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<td>1.220</td>
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<td>0.998</td>
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<td>55</td>
<td>5.970</td>
<td>1.075</td>
<td>0.583</td>
<td>0.008</td>
<td>0.912</td>
<td>0.018</td>
<td>0.997</td>
<td>0.364</td>
</tr>
</tbody>
</table>

The amount of acid that interacted with the standard slurries (Slurry-Standard-SW and Slurry-Standard-W) showed higher values than for modified slurries (Slurry-Mod-SW and Slurry-Mod-W). Moreover, the increase in temperature favors the interaction of the slurries with the acid and with and with greater proportion for the slurries without the Epoxy-EDA polymer. Therefore, it can be predicted that standard slurries have higher affinity for the mud-acid solution than the modified slurries.

The analysis of the kinetic parameters indicates that the acid attack is slower in the additives slurries, since it present smaller kinetic constants (k), when compared with the standard slurries. Temperature is an important factor to acid attack onto cement slurries. Normally, chemical reactions occurring at high temperatures give more energy to the system and increases the rate of reaction, causing more collisions between particles, as explained by collision theory. However, the main reason for temperature to increase the reaction rate is that more particles in the collision will have the required activation energy resulting in successful collisions. By analyzing some known effects, such as increase of both reaction rate and solubility of the calcium salts, it can be expected that temperature rise will increase the acid aggressiveness to the cement [55]. In the present study, it was observed that the mud-acid/cement interaction processes are temperature dependent. As temperature increases, the kinetic constants have high values, suggesting fast reaction rates and, thus, more intense acid attacks. The standard slurries have higher values of k than the modified slurries, suggesting that the standard slurries suffered greater aggression from the mud-acid solution.

The values of the kinetic parameter n can provide information concerning the mechanistic features of interaction phenomena at the solid/solution interfaces. Typically, values of n above 1.0 suggest interaction mechanisms occurred on the surface of the slurry. On the other hand, values of this parameter smaller than unity suggest that the mud-acid diffusion process govern these mud-acid/cement interactions [20]. In general, the values of n are close to unity. However, for the standard slurries, the values are lower than for the modified slurries, suggesting diffusion as the predominant mechanism of the attack for the control slurries. At the same time, it suggests that the slurries added with the Epoxy-EDA polymer should make it difficult to diffuse mud-acid into these slurries.

According to the kinetic data, the modified slurries show less interaction with the acid solution, when the reaction occurs at a slower speed. In addition, there is an apparent difficulty of diffusion of the H+ ions into the modified-slurries, suggesting also that the inclusion of such epoxy-based polymeric net offers some degree of protection in cement matrix against acid attack.
3.3 Evaluation of the mud-acid interaction with cement slurry by multivariate analysis and modeling

Factorial design is a fast and effective method for understanding and optimizing products and processes. The 2^4 factorial design is one of the most utilized in the research of effects of different factors on a specific response, where k is the number of factors analyzed in two levels [55].

The full 2^4 factorial design was performed to evaluate the importance of the parameters: cement slurry type, type of water used for preparation of the slurries, concentration of the mud-acid solution and the temperature on the mud-acid interaction onto the slurries. The responses evaluated were the amount of acid that interacts with the slurry (Q) and the kinetic constant (k), obtained from the kinetic modeling performed with the experimental data. These parameters were selected due to their importance in describing the interactions that occurred in the solid/solution interface [34].

The experimental design was performed using two levels for each selected factor, which were coded as (+) for the higher level and (-) for the lower level, as presented in Table 4. The effect of each factor shows the variation in the response caused by the change from the lower level factor (-1) to the higher level factor (+).

Table 4: Factors and levels used in the 2^4 factorial design study.

<table>
<thead>
<tr>
<th>FACTORS</th>
<th>LEVELS</th>
</tr>
</thead>
<tbody>
<tr>
<td>P = type of slurry</td>
<td>Standard</td>
</tr>
<tr>
<td>C= concentration of H^+ (mol.L^-1)</td>
<td>0.01</td>
</tr>
<tr>
<td>T = Temperature (ºC)</td>
<td>25</td>
</tr>
<tr>
<td>E= Environment for prepared of the slurries</td>
<td>Seawater</td>
</tr>
</tbody>
</table>

Table 5: Experimental and predicted results of the 2^4 factorial design.

<table>
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<tr>
<th>EXPERIMENT</th>
<th>P</th>
<th>C</th>
<th>T</th>
<th>E</th>
<th>Q.10^-4 EXPERIMENTAL (mol.g^-1)</th>
<th>Q.10^-4 PREDICTED (mol.g^-1)</th>
<th>k (h^-1) EXPERIMENTAL</th>
<th>k (h^-1) PREDICTED</th>
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<td>-1</td>
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<td>0.61</td>
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<td>-1</td>
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<td>4.20</td>
<td>4.19</td>
<td>0.223</td>
<td>0.227</td>
</tr>
<tr>
<td>13</td>
<td>-1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0.77</td>
<td>0.81</td>
<td>0.944</td>
<td>0.885</td>
</tr>
<tr>
<td>14</td>
<td>1</td>
<td>-1</td>
<td>1</td>
<td>1</td>
<td>0.63</td>
<td>0.61</td>
<td>0.705</td>
<td>0.701</td>
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<tr>
<td>15</td>
<td>-1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>8.20</td>
<td>8.19</td>
<td>0.644</td>
<td>0.639</td>
</tr>
<tr>
<td>16</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>5.70</td>
<td>5.71</td>
<td>0.529</td>
<td>0.523</td>
</tr>
</tbody>
</table>

Table 5 presents the factorial design 2^4 table. The efficiency of a given factorial design study is affected by the choice of the levels studied. In this study, it was observed that a significant difference between the levels occurred, resulting in a reliable evaluation of the factorial design responses. For Q, values ranged were between 5.20x10^-5 and 8.24x10^-4 mol. L^-1, and for k the range of values was: 0.22-0.944 h^-1. The main and interaction effects are presented in Table 6. The effects errors for each response evaluated were Q = 2.0x10^-6 mol. g^-1 and k = 3.0x10^-1 h^-1.

The factorial design has shown that the main effects for the kinetic responses are significant and influence the process of interaction between cement slurries and mud-acid solutions. The factors “Concentration of mud-acid solution”, “Temperature” and “Environment for prepared of the slurries” increase the main effects of Q. Increasing the concentration and temperature increases the quantitative extension of the mud-acid attack on the slurries. Increasing temperature, the mass transport acts as an important in the mud-acid interaction with the cement slurry. In general, interactions at the solid/solution interfaces increase with temperature,
because higher temperatures provide faster rates of diffusion of the solution species onto the solid surface [1].
In this way, a greater amount of acid interacts with the cement slurries.

Table 6: Effect values and their standard errors of the 2^4 factorial design.

<table>
<thead>
<tr>
<th>EFFECTS</th>
<th>Q (mol g^-1).10^4</th>
<th>k (h^-1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average</td>
<td>3.22 ± 0.01</td>
<td>0.560 ± 0.001</td>
</tr>
<tr>
<td>Principal</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P</td>
<td>-0.79 ± 0.02</td>
<td>-0.143 ± 0.003</td>
</tr>
<tr>
<td>C</td>
<td>5.17 ± 0.02</td>
<td>-0.204 ± 0.003</td>
</tr>
<tr>
<td>T</td>
<td>1.29 ± 0.02</td>
<td>0.288 ± 0.003</td>
</tr>
<tr>
<td>E</td>
<td>-0.08 ± 0.02</td>
<td>-0.013 ± 0.003</td>
</tr>
<tr>
<td>Interactions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>P-C</td>
<td>-0.69 ± 0.02</td>
<td>0.006 ± 0.003</td>
</tr>
<tr>
<td>P-T</td>
<td>-0.46 ± 0.02</td>
<td>-0.010 ± 0.003</td>
</tr>
<tr>
<td>P-E</td>
<td>-0.04 ± 0.02</td>
<td>-0.004 ± 0.003</td>
</tr>
<tr>
<td>C-T</td>
<td>1.15 ± 0.02</td>
<td>0.020 ± 0.003</td>
</tr>
<tr>
<td>C-E</td>
<td>-0.08 ± 0.02</td>
<td>-0.023 ± 0.003</td>
</tr>
<tr>
<td>T-E</td>
<td>-0.005 ± 0.02</td>
<td>-0.013 ± 0.003</td>
</tr>
<tr>
<td>P-C-T</td>
<td>-0.44 ± 0.02</td>
<td>0.021 ± 0.003</td>
</tr>
<tr>
<td>P-C-E</td>
<td>-0.03 ± 0.02</td>
<td>0.001 ± 0.003</td>
</tr>
<tr>
<td>P-T-E</td>
<td>-0.04 ± 0.02</td>
<td>-0.004 ± 0.003</td>
</tr>
<tr>
<td>C-T-E</td>
<td>0.01 ± 0.02</td>
<td>-0.011 ± 0.003</td>
</tr>
<tr>
<td>P-C-T-E</td>
<td>-0.02 ± 0.02</td>
<td>0.008 ± 0.003</td>
</tr>
</tbody>
</table>

Water is an important component of cement slurry, contributing actively to the hydration of the cement and its workability when fresh. All compounds present in the cement are anhydrous, but in contact with water it is hydrolyzed forming hydrated compounds. As the water participates in the formation of the cement slurry, the origin of this liquid must be evaluated during the process so that no problems occur in the properties in the cement slurry. However, in practice, water quality control is often neglected [15]. Thus, evaluating the quality of cement slurries in separate aqueous media becomes necessary and can provide economically and ecologically feasible responses for practical applications.

The aqueous medium, in which the slurries were prepared, seawater or distilled water, influences the process of interaction of cement slurries with mud-acid. The slurry prepared in seawater have a higher interaction with the acid than the slurry prepared in distilled water. The slurries prepared with seawater show greater affinity with the acid, corroborating with more corrosion in these slurries. By analyzing the slurry type factor, it is observed that its main effect decreased, indicating that the interaction of the acid species is lower for the Slurry-MOD than the standard slurries.

The temperature factor increased the main effect of k. The interaction of the acid mud with the cement slurries occurs more rapidly at higher temperatures. At lower temperatures, the interaction of mud-acid is driven by external mass transport, and the interaction rate constants (k) are lower than the interaction rate constants at high temperatures. When temperature increases, the thickness of the boundary layer around the slurry surface decreases and the resistance to mass transport of the acid species in solution increases [32].

The factors cement slurry cement, mud-acid concentration and Environment for prepared of the slurries decreased the main effect of k. At lower concentrations, the acid mud is consumed rapidly and a reaction enters into equilibrium. However, the acidic corrosion intensity increases with increasing mud-acid concentration. The resistance of the surface layer also depends on concentration of the acid solution. Increasing acid concentration may result in a variation in the composition of the corroded surface layer of the slurry, due to increased decalcification or dissolution of corrosion products [56].

Despite the higher interaction of the acid with the slurries prepared in seawater, the speed of this reaction was slower in relation to the slurry prepared with distilled water. The reaction rate for the Epoxy-EDA slurries was lower than for the standard slurry, suggesting that the diffusion of the acid occurred more slowly in the modified slurries compared to the standard slurries.

The effects interactions most significant of Q were Cement Slurry Type-Mud-acid Concentration, Temperature- Mud-acid Concentration, and Temperature- Environment for prepared of the slurries. Only two
interactions are not significant with respect to effects errors, Cement Slurry Type-Temperature and Cement Slurry Type-Mud-acid Concentration-Temperature. In the interactions where the Cement Slurry Type factor is present, the interaction decreases, suggesting that by changing the factors, the Slurry-MOD has less interaction with the acid than the standard slurry.

The interactions of the most significant k effects were: Mud-acid concentration - Environment for prepared of the slurries, Cement Slurry Type -Temperature, Mud-acid Concentration -Temperature. Some interactions were not considered statistically significant according to their effects errors.

Quantitative polynomial models for Q and k can be written using the statically significant effects, from data of Table 5:

\[
Q \times 10^4 = 3.22 - 0.24x_1 + 2.58x_2 + 0.65x_3 - 0.04x_4 - 0.35x_1x_2 - 0.23x_1x_3 - 0.02x_1x_4 + 0.58x_2x_3 - 0.04x_3x_4 - 0.22x_1x_2x_3 - 0.02x_1x_3x_4
\]

\[
k = 0.560 - 0.07x_1 - 0.1x_2 + 0.14x_3 - 0.007x_4 + 0.003x_1x_2 - 0.005x_1x_3 + 0.01x_2x_3 - 0.011x_3x_4 - 0.006x_4x_4 + 0.01x_1x_2x_3 - 0.005x_2x_3x_4 + 0.004x_1x_2x_4
\]

where \(x_1, x_2, x_3\) and \(x_4\) are codified (±1) values of slurry, concentration of mud-acid, temperature and environment for prepared of the slurries, respectively.

The experimental and predicted values of Q and k are also shown in Table 5. Good correlations are observed relating the observed and predicted values of Q and k.

4. CONCLUSIONS

In this study, an epoxy-ethylenediamine cement slurries was synthesized in two aqueous environments: distilled water and seawater. The characterization has pointed out significant compositional differences of the slurries. It was found lower amounts of portlandite in the polymer-modified slurries. The crystalline patterns of such slurries were preserved, before and after mud-acid interactions. On the other hand, the standard slurries have presented significant modifications after contact with mud-acid solution. It is noted that the standard slurries decreased drastically their crystalline features after mud-acid interaction. The analysis of the textural properties of the cement slurries indicated that the control slurries have greater porosity before and after the acid attack than the modified slurries, contributing to a lower chemical and mechanical resistance. Mechanical tests have shown that control slurries have less mechanical resistance than modified slurries.

The kinetic data of the interaction of the additive slurries with acidic species of mud-acid in solution were well fitted to an order-variable exponential kinetic model. The analysis of the kinetic modeling and characterization suggests that surface reactions with preservation of the chemical structures constitute the main mechanistic interactions at the interface of polymer-modified cement slurries, even after long-term contact with mud-acid.

Additional testing procedures must be performed to validate the use of the Epoxy-EDA-modified cement slurries in oil well cementing operations. However, the characterization and the factorial design results have suggested that this new cement slurry presents good potential to be used in innovative environmentally-friendly oil well procedures.

5. BIBLIOGRAPHY


ORCID
Danilo Oliveira Santos http://orcid.org/0000-0003-3306-429X
Ivory Marcos Gomes dos Santos https://orcid.org/0000-0001-5699-0080
Joenesson Filip Santos Ribeiro https://orcid.org/0000-0001-9188-5585
Eunice Fragoso Silva Vieira https://orcid.org/0000-0003-2599-3208
Gwenn Le Saoût https://orcid.org/0000-0002-2761-0584
Antonio Reinaldo Cestari https://orcid.org/0000-0002-7594-3279