Evaluation of Mechanical compressive strength of cementitious matrix with 12% of IER formulated by modified polymer (NEPS) at different percentages

Atiqa Bekhta1, Rachid Hsissou1,2* & Ahmed Elharfi1

During this paper, we improved the compressive strength of cementitious matrix based on ion exchanging resin (IER) at 12% and formulated by the modified novolac epoxy polymer surfactant (NEPS) at various percentages (0, 1, 2, 3, 4 and 5%). The results show that the introduction of 1% and 2% of NEPS in the cementitious matrix with 12% of IER increases the compressive strength compared to that of the basic matrix (from 7 to 90 days). However, the formulations 3, 4 and 5% show the compressive strength is less than that of the basic matrix (from 28 to 90 days).

Epoxy polymers cover various industrial areas such as: electronics, coating, inhibition, space construction and radioactive waste1−5. Epoxy polymers have many properties, including viscosimetric, thermal, morphological, rheological and mechanical compressive of strength6−9. At this time, the most widely employed process for the synthesis of epoxy polymers is the condensation of epichlorohydrin on structures having at least two mobile hydrogen atoms such as diamines, diacids and polyphenols. Also, the oxidation of polyunsaturated compounds and/or the condensation of glycridol with halogenated compounds10,11. Novolac epoxy polymers are obtained by the condensation of phenol with formaldehyde in an acid medium12,13.

The objective of this paper is to improve the chemical compressive of strength of the cementitious matrix based on ion exchanging resin (IER) at 12% formulated by novolac epoxy polymer surfactant (NEPS) modified at various percentages (1, 2, 3, 4 and 5%)14−16. We have studied the properties of mechanical of compressive strength, varying the polymer amounts incorporated in cementitious matrix formulations. This gave us the reflection to introduce the modified novolac epoxy polymer in surfactant form into the cementitious matrix. Besides, the results obtained show an increase in mechanical compressive of strength after 7, 14, 28 and 90 days of confinement compared to the basic matrix.

Material and Methods

Ion exchange resin (IER). Ion exchange resin is a crosslinked macromolecular water insoluble matrix which, upon contact with a solution, can exchange the ions it contains with other ions of the same sign from the solution used in water purification of the reactor vessel Mark TRIGA II16. Their physicochemical properties are shown in Table 1.

Novolac epoxy polymer (NEP). Epoxyes are excellent matrices of high performance polymers. The latter are synthesized by polycondensation reaction and are used in several fields such as conditioning of radioactive

1Laboratory of Agricultural Resources, Polymers and Process Engineering, Team of Organic Chemistry and Polymers, Department of Chemistry, Faculty of Science, University Ibn Tofail, BP 133, 14000, Kenitra, Morocco. 2Team of Innovative Materials and Mechanical Manufacturing Process, ENSAM, University Moulay Ismail, B.P. 15290, Al Mansour, Meknes, Morocco. *email: r.hsissou@gmail.com
waste reinforced concrete. The NEP is prepared by condensation of epichlorohydrin with polycresol (hydroxy novolac) polymer in an alkaline medium (Scheme 1).

**Cement.** The used CPJ 45 cement has technical characteristics which are conformed to the Moroccan standard NM 10.1.004. The CPJ 45 is a composite Portland cement resulting from the milling of clinker (+70%), the complement of 100% of one or more secondary constituents such as fillers, Pozzolan or fly ash, gypsum to regulate the setting. This cement is also called a hydraulic binder because it has the property of hydrating and curing in the presence of water.

**Molds.** The used molds are cylindrical with a diameter of 5 cm and a height of 10 cm, which are illustrated in Fig. 1.

**Mixer.** The auto mortar mixer is a device which ensures the mixing of a great homogeneity while reducing the duration of mixing with 30 seconds of time and 285 rpm of speed of mixing.

**Press carver 4350 L.** The used press carver 4350 L is manual hydraulic which makes it possible to determine the compressive strength of the mortar from the force measured in view of the surface.

**Preparation of surfactant.** The synthesis of modified novolac epoxy polymer surfactant (NEPS) based on novolac epoxy polymer (NEP) was carried out in two steps. The first step consists of condensing of 0.004 mol of novolac epoxy polymer with 0.026 mol of acrylic chloride in the presence of Lewis acid (AlCl₃) by Friedel and Craft acylation reaction with magnetic stirring for 4 hours at 100 °C. Besides, in the second step, 3.014 mol of para-aminophenol were added to the previously product obtained (I) according to the first step by reaction of 1, 4-Mickael addition under magnetic stirring for 3 hours at 70 °C (Scheme 2). All the employed chemicals products were purchased from Aldrich Chemical Co.

**Calculation of compressive strength.** The strength is applied to the matrix by two cylindrical metal plates and the reading of the force is either pound or Pound per Square Inch (PSI). Moreover, we calculate the compressive strength (R) from the displayed force value which is expressed by using Eq. 1.

\[ R = \frac{F}{S} \]  

With R, F and S denote compressive strength (MPa), force applied (Pound) and surface of the test piece (cm²), respectively.

**Methods used**

**Fourier transform infrared spectroscopy.** The used FTIR spectrometer is a BRUKER Fourier transform spectroscopy. The light beam passes through the sample to a thickness of about 2 μm. The analysis is carried out between 4000 cm⁻¹ and 600 cm⁻¹.

**Nuclear magnetic resonance.** Analyses of Nuclear magnetic resonance (¹H NMR and ¹³C NMR) were obtained by using ADVANCE 300 Bruker like apparatus, and the product was solubilized in CDCl₃. The chemical shifts are expressed in ppm.

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| Physical and chemical properties of IER |  |
|---|---|
| **Skeleton** | Polystyrene crossed with DVB of gel type |
| **Functional groups** | R-SO₃⁻ |
| **Physical aspect** | Dark amber beads, translucent |
| **Ion shape on delivery** | H⁻ |
| **Moisture content** | 51–65% (H⁺ form) |
| **Maximum swelling** | Na⁺–H⁺: 5% |
| **Temperature limit** | 120 °C |
| **Limit of pH** | From 0 à 14 |
| **Apparent density** | Approximately 800 g/L |
| **Actual density** | 1.20 (H⁺ form) |
| **Total exchange capacity** | Min 1.7 eq/l (H⁺ form) |

**The granulometry of the resin**

| Less than 0.315 mm | 0.2 max |
| 0.4 < X < 1.0 mm | 80% min |
| Greater than 1.25 mm | 3% max |

**Table 1. Physical and chemical properties of IER.**
Scanning electron microscope. The scanning electron microscope was used to make photographic images. The observations were carried out on a JEOL-JEC-530 microscope. This technique is based on the use of a beam of electrons accelerated by a fixed potential that excites the surface of the sample.

Results and Discussions

Fourier transform infrared spectroscopy. The novolac epoxy polymer surfactant (NEPS) modified was characterized by Fourier transform infrared spectroscopy (FTIR) analysis (Fig. 2). The different bonds NEPS are grouped in Table 2.

Nuclear magnetic resonance. The proton and carbon NMR spectrum of the novolac epoxy polymer surfactant modified are shown in Figs. 3 and 4, respectively. The chemical shift (ppm) of proton and carbon of NEPS modified are given as follows.

$^1$H NMR (ppm): 2.38–2.63 (m, 2H, CH$_2$ of oxiranes); 2.78–3.25 (m, 4H, CH$_2$ of methylene); 3.04 (m, 2H, CH of oxiranes); 3.95 (s, 2H, CH$_2$ bonded to two benzene); 4.0 (m, 4H, C-NH); 4.20 (dd, 2H, C-H of oxiranes); 5.0 (s, 2H, C-OH) and 6–8 (d, 4H, C-H of benzene).
Table 2. Different bonds of NEPS.

| Band ν (cm⁻¹) | Attributions                                      |
|--------------|--------------------------------------------------|
| 3200         | Bond of O–H residual                             |
| 2920         | Bond stretching of CH₂                           |
| 1300-1500-1590 | Bond stretching of C–O                        |
| 1150         | Bond stretching of C–O aromatic ethers (Ph–O) and alcohols |
| 1100         | Bond of C–C aliphatic                           |
| 1020         | Bond of C–N                                     |
| 815          | Bending of CH₂ (epoxy)                          |
13C NMR (ppm): 26.48 (s, 1C, CH$_2$ bonded with two benzene); 36.4–39.2 (s, 4C, CH$_2$ of methylene); 44.2 (s, 2C, CH$_2$ of the oxirane); 50.7 (s, 2C, CH of oxirane); 69.5 (s, C of CH$_2$ bonded with ether); 114.6, 116.9 and 128.6 (s, C of benzene); 140.2 (s, C of benzene bound to NH); 146.5 159.1 (s, C of benzene bound to ether) and 200.0 (s, C of carbonyl).

Optimization of percentage of NEPS modified. The different formulations used for this study are 12% of IER, 67.92% of cement and 20.19% of water. In this study, we tried to introduce the novolac epoxy polymer surfactant (NEPS) modified into the cement matrix so as to improve the compressive strength. The used surfactant polymer is introduced in matrix at different percentages (1, 2, 3, 4 and 5%). Figure 5 shows the confinement matrix. The results of compressive strength for these 7, 14, 28 and 90 days matrix are shown in Table 3. According to these results, we concluded that the introduction of novolac epoxy polymer surfactant modified at various percentages (1, 2, 3, 4 and 5%) into the matrix increases the compressive strength with respect to the base matrix (Fig. 6). The matrix of 1% of NEPS modified has a superior compressive strength. Once again, we changed the configuration of the novolac epoxy polymer by modifying it in surfactant form to further improve
the compressive strength in the conditioning matrix. Moreover, the compressive strength increases with time for formulations of 1 to 5% novolac epoxy polymer surfactant and for base formulation. Furthermore, the compressive strength of the matrices with the novolac epoxy polymer surfactant of this test is higher than that of the base matrix up to 28 days, from 28 days to 90 days, whereas the test’s matrix 3, 4 and 5% is less than that of the basic matrix14. Besides, the evaluation of these results shows that the introduction of 1% and 2% of NEPS into the IER conditioning matrix increases the compressive strength with respect to the base matrix16. They, thus, make it possible to solubilize two immiscible phases. For this reason, we have given a good homogeneity of the conditioned cementitious matrix, as well as good dispersion.

Table 3. Compressive strength (CS) of matrix based on NEPS modified (all in MPa).

| Times (d) | CS (0% NEPS) | CS (1% NEPS) | CS (2% NEPS) | CS (3% NEPS) | CS (4% NEPS) | CS (5% NEPS) |
|----------|--------------|--------------|--------------|--------------|--------------|--------------|
| 7        | 8.06         | 12.94        | 17.0625      | 11.25        | 9.375        | 9.187        |
| 14       | 8.43         | 16.31        | 17.44        | 11.63        | 10.875       | 10.69        |
| 28       | 11.06        | 19.69        | 19.31        | 9.56         | 10.31        | 10.78        |
| 90       | 10.87        | 17.81        | 17.34        | 9.38         | 9.187        | 9.28         |

Figure 6. Variation of compressive strength according to time.

Figure 7. Micrographs of different formulations prepared (0, 1, 2, 3, 4 and 5%).
Scanning electron microscopy. The dispersion of ions exchanging resin (IER) in the cementitious matrix formulated by novolac epoxy polymer surfactant at various percentages is presented in Fig. 7. The different formulations with addition of 0 to 5% of NEPS modified are analyzed by the scanning electron microscopy (SEM). According to the SEM micrographs observations, the cementitious matrix formulated by NEPS clearly show the spherical IER loads on the analyzed surfaces9,22. The addition of 1% NEPS exhibits better dispersion, this confirms the higher compressive strength.

Conclusion

Several research studies have been conducted so as to determine which formulation has better compressibility resistance than the IER conditioning base formulation. Previous studies have evaluated the impact of the novolac epoxy polymer surfactant (NEPS) in different physical states on the formulation. The objective of this study was to improve the compressive strength of the containment matrix by setting the percentage of IER at 12%, the percentage of water at 20.19% and cement at 67.92%, and the introduction of the novolac epoxy polymer surfactant (NEPS) to different percentages (1, 2, 3, 4 and 5%). Besides, the results obtained in this study showed an increase in the compressive strength after 7, 14, 28 and 90 days of confinement with respect to the base formulation. In addition, the introduction of novolac epoxy polymer surfactant into formulation allowed an improvement in the compressive strength of 1% and 2% matrix of the NEPS and good homogeneity of the conditioned cementitious matrix, as well as good dispersion of IER in our formulations on the other hand.

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Atiqa Bekhta prepared all figures, Rachid Hsissou written and interpreted all experimental works and Ahmed Elharfi designed the work and was part of the manuscript writ-up.

Competing interests
The authors declare no competing interests.

Additional information
Correspondence and requests for materials should be addressed to R.H.

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