An Analysis of Crystalline Admixtures in Terms of Their Influence on the Resistance of Cementitious Composites to Aggressive Environments

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Received: 05 June 2019, Accepted: 06 November 2020, Published online: 23 November 2020

Abstract
This paper describes the influence of crystalline admixtures on the chemical resistance of cement composites exposed to aggressive environments. The effect of the crystalline admixtures was determined by a series of physical-mechanical and innovative physical-chemical methods. Specifically, this concerned the measurement of flexural strength, compressive strength, determination of the dynamic modulus of elasticity by the ultrasonic pulse velocity test, and an analysis of the internal structure by mercury intrusion porosimetry and x-ray computed tomography. Physical-chemical analyses were also performed; namely an x-ray diffraction analysis to determine the mineralogical composition and electron microscopy to examine the microstructure. The use of non-destructive testing methods (ultrasonic pulse velocity test and computed tomography) made it possible to compare the properties of the same specimens for 16 months. The specimens were stored in reference laboratory conditions, a sodium sulphate solution and an ammonium chloride solution. The physical-mechanical tests and physical-chemical analyses clearly showed the benefit that crystalline admixtures have for the resistance of cementitious composites attacked by chemically aggressive solutions without affecting the fresh-mixture rheology or decreasing the strength of the composites.

Keywords
cementitious composites, crystalline admixtures, aggressive environment, resistance, modern diagnostic methods

1 Introduction
Contemporary civil engineering demands materials of excellent properties, which means not only good strength but also durability under all usage conditions.

One of the useful principles that can be used to improve the resistance of cement composites to adverse environments involves using suitable admixtures to prevent the ingress of aggressive agents through the capillary pore structure.

In recent years, researchers have been focusing on self-healing in ordinary concrete [1], self-healing engineered with fibber reinforcement [2–4], bacteria-based self-healing concrete [5], super absorbent polymers [6], self-healing of mortars using tubular capsules made from glass and ceramics filled with a healing agent [7], self-healing effect of microcapsules with an oil core and a silica gel shell [8], and other admixtures such as flyash [9], nanographite [10–12], marble powder [12–16] and brick powder [14–16]. In the latter case, the self-healing occurs mainly by the filling of cracks, swelling and expansion, and improved hydration and re-crystallization [1].

The self-healing mechanism requires a sufficient amount of water to be present, especially when chemical agents were added to promote the growth of crystals inside the cracks.

One of the promising new technologies for protecting cementitious composites against aggressive environments is the use of crystalline admixtures, which ACI TC 212 [17] classifies as a type of permeability-reducing admixtures (PRA). Crystalline admixtures (CA) are hydrophilic, i.e., they react easily with water, in contrast to water-repellent or hydrophobic products [18]. The crystalline formations produced by CA become a permanent part of the cement matrix [6, 17].
Their behavior is still not fully understood. In fact, the ACI TC 212 report [17] states that the concrete compounds that react with CA are tricalcium silicates, although other authors [18] state calcium hydroxide as the reactive. The process, according to [17], following equation (Eq. (1)), where a crystalline promoter \( \text{M}_x \text{R}_x \) reacts with tricalcium silicates and water to produce modified calcium silicates hydrates and a pore-blocking precipitate \( \text{M}_x \text{CaR}_x - (\text{H}_2\text{O})_x \).

\[
3\text{CaO} - \text{SiO}_2 + \text{M}_x \text{R}_x + \text{H}_2\text{O} \rightarrow \text{Ca}_x \text{SiO}_3 \text{R} - (\text{H}_2\text{O})_x + \text{M}_x \text{CaR}_x - (\text{H}_2\text{O})_x
\]

(1)

The research described herein focused on assessing the effect of this type of CA.

2 Materials and experimental procedures
2.1 Specimen preparation and exposure to aggressive environments
This particular experiment analyzed the chemical resistance delivered by CA using fine-grained cement composites (aggregate fraction according to CSN EN 196-1 [19]).

It tested CA-modified materials as well as materials modified by a combination of the admixtures and dispersed reinforcement.

One of the important points of the research was a comparison of the modified materials with unmodified ones, i.e. with only a pure cement matrix (identified as I through III). Table 1 shows the composition of the test mixtures.

Water was added so as to keep the workability more or less constant, following the determination of consistency of fresh mortar according to CSN EN 1015-3 [20] was 160±10 mm.

Table 2 shows the values of consistency determined by the flow table test.

The tests have shown that the workability of the fresh mixtures is more affected by the presence of dispersed reinforcement rather than the crystalline admixture, which has only a minor influence on this parameter.

The mixtures contained the following raw materials:

Cement CEM I 42.5 R was used as binder, sand was added in accordance with CSN EN 196-1 [19] and the CA was XYPEX Admix (XYPEX CHEMICAL CORPORATION, Richmond B.C., Canada). The second batch of specimens was made with polypropylene fibres Fibrin 3/15 (KrampeHarex CZ, s.r.o., Czech Republic).

Following a procedure described in CSN EN 196-1 [19] the mixtures were cast into prisms of 40 × 40 × 160 mm.

The specimens were stored for 28 days at a temperature of 23°C and relative humidity of 95%. At the age of 28 days they were placed in aggressive solutions; see Table 3 for details. Some of the specimens remained in laboratory conditions throughout the experiment and served for reference.

The concentration of the solutions was monitored throughout the experiment and kept at a constant level.

2.2 Testing procedures and analysis
At selected time intervals (exposure of 3, 6, 12 and 15 months), some of the specimens were removed from their environments and subjected to physical-mechanical and physical-chemical tests:

- Determination of flexural strength \( (R_f) \) and compressive strength \( (R_c) \) according to [19],
- Determination of the dynamic modulus of elasticity; it was measured every month by the ultrasonic pulse velocity test. The ultrasonic pulse velocity test is a non-destructive method used mainly in concrete testing and is standardized by ISO 1920-7 [21], ASTM C597 [22], CSN EN 12504-4 [23], CSN 731371 [24], CSN 731380 [25]. The parameters measured by the ultrasonic pulse velocity test can be used to determine physical-mechanical properties (dynamic modulus of elasticity, strength estimation) as well as changes in the internal structure due to

| Mix. ID | Cement [kg] | Sand [kg] | CA [kg] | PP Fibres [kg] | Water [L] |
|---------|-------------|-----------|---------|----------------|-----------|
| I       | 512         | 1535      | 0       | 0              | 256       |
| II      | 511         | 1533      | 3       | 0              | 256       |
| III     | 510         | 1531      | 8       | 0              | 255       |
| IV      | 511         | 1533      | 0       | 1              | 262       |
| V       | 511         | 1532      | 3       | 1              | 264       |
| VI      | 510         | 1539      | 8       | 1              | 263       |

| Mix. ID | Flow value [mm] |
|---------|-----------------|
| I       | 158             |
| II      | 159             |
| III     | 158             |
| IV      | 155             |
| V       | 153             |
| VI      | 157             |

| Environment ID | Specification |
|---------------|---------------|
| Laboratory conditions - reference | temperature of 23°C and relative humidity of 95% |
| Sodium sulphate solution | concentration of 36 000 mg of SO\(_4^{2-}\) |
| Ammonium chloride solution | concentration of 3 000 mg of NH\(_4^+\) |
The internal structure was analyzed using mercury intrusion porosimetry and x-ray computed tomography (CT) using the method described in [31]. Various other authors have used the ultrasonic pulse velocity test to assess changes in the internal structure of concrete and other building materials, e.g. [26–30].

The research described here used this method (as described in CSN 731380 [25]) to observe changes in the internal structure of secondary crystallization-modified cement composites exposed to aggressive environments. These changes were assessed based on changes in the relative dynamic modulus of elasticity depending on time of exposure.

Test equipment – an ultrasonic tester with an accuracy of 0.1µs and natural transducer frequency of 150 kHz. Measurement procedure and evaluation - the specimens with the dimensions of 40 × 40 × 160 mm were tested at 30-day intervals by direct transmission in the longitudinal direction. The measurement outcome is the transit time T. Relative dynamic modulus of elasticity \( RDM_{UPPT,n} \) (CSN 731380 [25]) was calculated using Eq. (2):

\[
RDM_{UPPT,n} = \left( \frac{t_{S,0}}{t_{S,n}} \right)^2 . 100 ,
\]

where:

\( RDM_{UPPT,n} \) - relative dynamic modulus of elasticity calculated from the ultrasonic pulse time after \( n \) months of exposure to an aggressive medium [%].

\( t_{S,0} \) - initial transit time [µs];

\( t_{S,n} \) - ultrasonic pulse transit time after \( n \) months of exposure to an aggressive medium (\( n = 1 \) through 14 months) [µs];

- The internal structure was analyzed using mercury intrusion porosimetry and x-ray computed tomography (CT) using the method described in [31]. An x-ray (CT) image analysis showed to be a powerful technique to study pore-related characteristics and create simulations of cementitious materials [32–34].

- Physical-chemical analyses involved the determination of mineralogical composition using x-ray diffraction (XRD) and differential thermal analysis (DTA). The microstructure was also examined by scanning electron microscopy. These tests were performed following a method described in [35].

The use of non-destructive testing methods (ultrasonic pulse velocity test and computed tomography) made it possible to make tests on the same specimens. In terms of determining the durability of building materials, this approach is fairly unique and very effective.

### 3 Experiment results

#### 3.1 Physical-mechanical parameters

Table 4 shows the results of flexural and compressive strength tests for specimens stored in laboratory conditions and Tables 5 and 6 show the strength of specimens attacked by the aggressive media (sulphates and chlorides). The values in the tables are averages made from test sets, each consisting of six specimens.

\( a) \) Aggressive sulphate environment

Graphs 1 and 2 show changes in flexural and compressive strength in dependence on the length of exposure to the aggressive environment and type of material. The strength

| Table 4 Strength – laboratory conditions |
|--------------------------------------------------|-------------------|----------|----------|----------|----------|----------|----------|
| Mix. ID | | 1 month | 4 months | 7 months | 13 months | 16 months |
| | \( R_f [\text{MPa}] \) | \( R_c [\text{MPa}] \) | \( R_f [\text{MPa}] \) | \( R_c [\text{MPa}] \) | \( R_f [\text{MPa}] \) | \( R_c [\text{MPa}] \) |
| I | 9.8 | 35.2 | 9.7 | 55.5 | 9.4 | 53.3 |
| II | 11.2 | 54.1 | 10.7 | 56.2 | 11.1 | 53.5 |
| III | 11.4 | 54.4 | 11.1 | 56.4 | 11.5 | 53.9 |
| IV | 12.0 | 54.7 | 11.4 | 56.7 | 11.8 | 54.1 |
| V | 12.1 | 54.3 | 11.6 | 56.8 | 12.1 | 54.3 |
| VI | 12.3 | 55.1 | 11.9 | 55.1 | 12.4 | 55.3 |
| | \( R_f [\text{MPa}] \) | \( R_c [\text{MPa}] \) | \( R_f [\text{MPa}] \) | \( R_c [\text{MPa}] \) | \( R_f [\text{MPa}] \) | \( R_c [\text{MPa}] \) |
| I | 52.6 | 52.7 | 53.7 | 54.1 | 54.2 | 54.7 |
| II | 53.9 | 54.1 | 54.7 | 55.1 | 54.1 | 54.3 |
| III | 54.1 | 54.3 | 54.7 | 55.1 | 54.2 | 54.7 |
| IV | 54.7 | 55.1 | 54.2 | 55.1 | 54.2 | 54.7 |

Note: Table 4 shows values of flexural strength \( (R_f) \) and compressive strength \( (R_c) \), which served as reference during evaluation. Specimens were placed in the aggressive environments at an age of 1 month; the exposed specimens were thus compared with references of 1 month + exposure time.
changes are expressed as percentage, where 100 % are reference values measured on specimens of the corresponding age (i.e. specimens ageing in laboratory conditions) (Figs. 1–2).

b) Aggressive chloride environment

The graphs below show how the solutions influence the strength of the materials. The effect is written in percentages, where 100 % represent values measured on the reference specimens of an adequate age (i.e. specimens ageing in laboratory conditions) (Figs. 3–4).

3.2 Analysis of internal structure

a) Ultrasonic pulse velocity test

Changes in the internal structure of materials exposed to the aggressive environments (sulphate and chloride solutions) were determined as a difference in the relative dynamic modulus of elasticity measured by the ultrasonic pulse velocity test.

| Mix. ID | Exposure time | 3 months | 6 months | 12 months | 15 months |
|---------|---------------|----------|----------|-----------|-----------|
| I       | $R_f$ [MPa]   | 12.6     | 12.1     | 11.0      | 8.4       |
|         | $R_c$ [MPa]   | 54.1     | 52.6     | 49.8      | 47.5      |
| II      | $R_f$ [MPa]   | 12.5     | 12.1     | 11.8      | 10.0      |
|         | $R_c$ [MPa]   | 56.2     | 54.3     | 50.1      | 49.4      |
| III     | $R_f$ [MPa]   | 12.7     | 12.4     | 12.1      | 10.5      |
|         | $R_c$ [MPa]   | 53.5     | 51.1     | 51.8      | 51.0      |
| IV      | $R_f$ [MPa]   | 12.9     | 12.6     | 10.1      | 9.5       |
|         | $R_c$ [MPa]   | 51.5     | 51.7     | 50.9      | 48.7      |
| V       | $R_f$ [MPa]   | 12.4     | 12.8     | 11.9      | 11.6      |
|         | $R_c$ [MPa]   | 51.2     | 50.9     | 50.2      | 49.3      |
| VI      | $R_f$ [MPa]   | 12.3     | 12.6     | 11.8      | 11.4      |
|         | $R_c$ [MPa]   | 52.7     | 52.1     | 50.4      | 50.2      |

| Mix. ID | Exposure time | 3 months | 6 months | 12 months | 15 months |
|---------|---------------|----------|----------|-----------|-----------|
| I       | $R_f$ [MPa]   | 11.2     | 12.1     | 11.0      | 9.2       |
|         | $R_c$ [MPa]   | 55.2     | 53.4     | 52.1      | 51.9      |
| II      | $R_f$ [MPa]   | 10.9     | 11.3     | 11.0      | 11.0      |
|         | $R_c$ [MPa]   | 57.4     | 55.6     | 57.8      | 54.9      |
| III     | $R_f$ [MPa]   | 10.9     | 11.7     | 11.4      | 11.2      |
|         | $R_c$ [MPa]   | 52.9     | 52.1     | 52.3      | 52.1      |
| IV      | $R_f$ [MPa]   | 10.8     | 11.2     | 11.4      | 10.9      |
|         | $R_c$ [MPa]   | 52.3     | 53.1     | 50.5      | 50.2      |
| V       | $R_f$ [MPa]   | 11.5     | 11.7     | 10.9      | 10.4      |
|         | $R_c$ [MPa]   | 53.4     | 52.5     | 51.6      | 50.8      |
| VI      | $R_f$ [MPa]   | 11.4     | 12.1     | 11.8      | 11.7      |
|         | $R_c$ [MPa]   | 52.9     | 53.1     | 51.4      | 51.0      |
The progress of the relative dynamic modulus of elasticity is plotted in Fig. 5 (sulphates) and Fig. 6 (chlorides).

b) Mercury intrusion porosimetry and computed tomography

In order to assess the effect of CA and its ability to interact with cement hydration products the internal structure of the composites, i.e. their porosity, was examined.

New specimens were made from a cement paste. Reference specimens contained no admixtures, while the test specimens contained a CA at an amount of 1.5 % of cement mass. They had the shape of a cylinder with the dimensions of 80 × 100 mm. They were left to age for 28 days in laboratory conditions. After this time, they were monitored at 28-day intervals to ascertain whether and to what extent the CA affects the capillary pore structure of the material being tested. This determination was carried out throughout the entire experiment, i.e. for 15 months.

The results of the measurements show that as the CA interacts with the hydration products, the pores are slowly eliminated; especially ones from 300 to 400 μm in size. This finding was proved both by mercury intrusion porosimetry and by computed tomography.

For illustration, see the CT images of mixture III below (i.e. mixture with 1.5 % of CA). These images were taken on the same specimen; Fig. 7 was taken at the age of 28 days and Fig. 8 shows the capillary pore structure at the age of 15 months.
3.3 Physical-chemical analysis

Other analytical methods used during the experiment were XRD, DTA, and an examination by a scanning electron microscope.

Similarly to the previous case, the specimens were left to age in laboratory conditions for 28 days and then exposed to the aggressive environments. The three analyses were then performed at 28-day intervals.

Special attention was paid to the identification of new products forming as a result of the action of the aggressive environments or other negative changes taking place in the material matrix. An important aspect was the comparison between the results of the microstructure analyses performed on specimens exposed to the aggressive environments and reference specimens.

Given the fact that the strength tests registered the most marked changes after exposure to the sulphate solution, the Table 7 shows the results of an XRD analysis performed on specimens exposed to this solution.

Another of the analyses was differential thermal analysis. Its goal was mainly to supplement and expand the findings obtained by XRD.

A key finding delivered by this analysis is the clear decrease in portlandite detected in mixtures containing no CA and exposed to a sulphate solution. On the other hand, CA-modified mixtures the portlandite decrease was significantly smaller. The graph below plots the change in portlandite content brought about by the sulphate solution in dependence on mixture composition and exposure time (Fig. 9).

Another method of analysis used herein was scanning electron microscopy, which focused on a detailed examination of typical features identified in the materials, especially in those exposed to chemical attack.

Among other things, this analysis showed that corrosion products form in the capillary pore structure of the unmodified composites.

In the modified materials, the occurrence of these products is significantly limited by the effect of the CA, as are other defects, such as cracks, even after long-term exposure to aggressive environments. The figures below illustrate these findings (Figs. 10–11).

4 Discussions

The results of the physical-mechanical analyses can be summarized as follows:

- It was proved that the addition of a CA has virtually no effect on the fresh mixture, not even at 1.5 % of cement mass. The workability was slightly influenced by the use of polypropylene fibers.

| Mix. ID | Laboratory conditions (reference) | Time of sulphate exposure |
|---------|-----------------------------------|---------------------------|
|         |                                   | 3 months | 6 months | 12 months | 15 months |
| I       | K, Po, CSH, Q                     | K, Po, CSH, Q | K, CSH, A, Q | K, CSH, A, E, MS, Q | K, E, MS, S, Q |
| II      | K, Po, CSH, trace C, Q            | K, Po, CSH, Q | K, Po, CSH, Q | K, CSH, A, E, trace E, Q | K, CSH, A, trace E, Q |
| III     | K, Po, CSH, A, Q                  | K, Po, CSH, A, Q | K, Po, CSH, A, Q | K, Po, CSH, A, Q | K, Po, CSH, A, Q |
| IV      | K, Po, CSH, A, Q                  | K, Po, CSH, A, Q | K, Po, CSH, A, Q | K, CSH, E, S, Q | K, CSH, E, S, Q |
| V       | K, Po, CSH, Q                     | K, Po, CSH, Q | K, Po, CSH, Q | K, Po, CSH, Q | K, Po, CSH, Q |
| VI      | K, Po, CSH, Q                     | K, Po, CSH, Q | K, Po, CSH, Q | K, Po, CSH, Q | K, Po, CSH, Q |

Note: Table 6 uses the following abbreviations:

K - calcite; Po - portlandite; CSH - calcium hydro silicate gels; C - carbonate complex; A - aragonite; E - ettringite, MS - monosulphate; S - gypsum; Q - β quartz.

![Fig. 9 Determination of the amount of portlandite in mixtures exposed to a sulphate solution](image-url)

- The effect of CA on the composites' capillary porosity was tested following a methodology described in ČSN 73 1316 [35] Determination of moisture content, absorptivity and capillarity of concrete. A series of tests has shown that the addition of a CA improves sorption properties. This statement can be illustrated e.g. by comparing the values of absorptivity which reached 7.3 % in mixture I (no CA content) as opposed to mixture III with 4.9 % (1.5 % CA content).
It was found that the effect of the aggressive environments is most strongly manifested in flexural strength, less so in compressive strength and Young’s modulus.

Strength changes in specimens exposed to sulphate attack were greater than in those exposed to ammonium chloride.

The positive effect of CA on strength was clearly demonstrated. This statement is documented by the fact that the 15-month exposure to a sulphate solution resulted in a 30.6% decrease in flexural strength in mixture I (unmodified) or 22.8% in mixture IV (unmodified, PP fibers). However, in mixtures that contained a CA the decrease did not exceed 14% and in modified mixtures with PP fibers it was even lower.

By extension, the benefit of CA was also confirmed in other strength characteristics (compressive strength, dynamic modulus of elasticity) in specimens exposed to the aggressive environments.

The analysis of the internal structure indicates the following:

- It was proved that exposure to the aggressive environments causes the relative dynamic modulus of elasticity to decrease, which indicates irreversible damage to the internal structure.

- For instance, after exposure to the sulphate solution, the unmodified specimens reached the inflection point after 8 months, whereas specimens with a CA reached it at approximately 11 months. A similar shift in inflection points between modified and unmodified materials was also found in the case of chloride attack. This is one of the indications of the benefit of CA in terms of their improving the chemical resistance of cement composites.

- The physical-chemical analyses proved that the CA can markedly limit undesirable changes taking place in the microstructure of materials exposed to aggressive environments. This is documented e.g. in the less pronounced decrease in portlandite content in specimens modified by the CA exposed to a sulphate solution brought about by a reduction in the number of corrosion products, etc.

- The process of interaction between the CA and cement matrix spans over a long period of time and it can thus be assumed that the experiment presented here was too short to capture the issue in full. Besides age, another important factor that influences changes in the character of the capillary pore structure are curing and ageing conditions, most notably humidity. This corresponds with research presented in [36].
5 Conclusions
The paper analyses the effect that crystalline admixtures have on fine-grained cement composites exposed to certain aggressive solutions.

The findings show that the use of dispersed reinforcement (fibres) affects workability to a greater degree than the addition of CA, which has only a minor influence.

Likewise, adding CA to the composites does not reduce their strength.

A series of physical-mechanical tests and physical-chemical analyses have clearly proved the positive effect they have on the chemical resistance of cement composites. This finding is in keeping with many previous investigations [37]. The main mechanism of this is the sealing of the capillary pore system which reduces the ingress of aggressive media into the internal structure of the composite.

In terms of the service life of steel-reinforced concrete structures the effect of CA can increase over the long-term horizon.

Acknowledgment
The paper was produced with the financial support of Czech Science Foundation, project No. 16-25472S "Secondary crystallization modified cement composites and their degradation dynamics".

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