

Effect of NaCl on the hydric and hygric dilation behaviour of lime-cement mortar

Lubelli, B.; Van Hees, R. P.J.; Huinink, H. P.

Publication date Document Version Final published version Published in Heron

Citation (APA)

Lubelli, B., Van Hees, R. P. J., & Huinink, H. P. (2006). Effect of NaCl on the hydric and hygric dilation behaviour of lime-cement mortar. *Heron*, *51*(1), 33-48.

Important note

To cite this publication, please use the final published version (if applicable). Please check the document version above.

Copyright
Other than for strictly personal use, it is not permitted to download, forward or distribute the text or part of it, without the consent of the author(s) and/or copyright holder(s), unless the work is under an open content license such as Creative Commons.

Takedown policy

Please contact us and provide details if you believe this document breaches copyrights. We will remove access to the work immediately and investigate your claim.

Effect of NaCl on the hydric and hygric dilation behaviour of lime-cement mortar

B. Lubelli
Delft University of Technology

R.P.J. van Hees

Delft University of Technology - TNO Built Environment and Geosciences

H.P. Huinink Eindhoven University of Technology

The mechanism of damage due to NaCl crystallization has not been clarified yet. Apart from crystallization pressure, other hypotheses have been proposed to explain the decay. Irreversible dilation during NaCl crystallization has been observed in a few cases but has never been studied in a systematic way. In this paper the effect of NaCl on the hydric and hygric behaviour of a lime/cement mortar is extensively studied. NaCl is shown to modify the hydric and hygric dilation behavior of the material. The material contaminated with NaCl shrinks during dissolution and dilates during crystallization of the salt. This dilation is irreversible and its amount is sufficient to cause damage after few dissolution/crystallization cycles. A similar behaviour has been also observed in the presence of NaNO₃ and KCl. If a crystallization inhibitor, modifying crystallization habits of the salt, is added to the NaCl, no irreversible dilation occurs. Outcomes of electron microscopy studies suggest that a relation exists between crystallization habit of the salt and dilation behaviour. Salts crystallizing as a layer adhering to the pore walls seem able to cause relevant expansion during drying of the specimen and crystallization of the salt. A similar effect is, in this theory, not possible when the same salt crystallizes without adhering to the material, as in the presence of the crystallization inhibitor.

Keywords: Salt crystallization, decay, dilation, hygric behaviour, hydric behaviour

1 Introduction

The damage mechanism due to NaCl crystallization has not been unequivocally explained yet. According to the most accepted theory, salt crystallization damage [Correns, 1949; Winckler and Singer, 1972; Scherer, 1999] occurs because of salt filling the pores and creating pressure on the pores' walls. This pressure depends on to the supersaturation ratio. In an equilibrium situation the supersaturation that can be reached depends on the pore radius: the smaller the pores the

higher the supersaturation. Following this theory, in order to produce crystallization pressures high enough to cause damage, NaCl should crystallize at high supersaturation filling pore sizes in the range of a few nanometers [Flatt, 2002; Rijniers et al, 2005]. Some objections have been raised to these theoretical conclusions: NaCl has a low tendency to supersaturate, therefore would hardly produce high crystallization pressures [Pel et al, 2002]. Besides, most traditional building materials do not have pores in the nanometer range but show, nevertheless, serious damage in the presence of NaCl.

The reported discussion suggests that damage due to NaCl, may not only be related to crystallization pressure as described in literature starting with Correns' work, but involves other mechanisms. The lack of systematic experimental investigations of the phenomenon of irreversible dilation in the presence of salts makes difficult to evaluate the importance of this phenomenon in the development of sodium chloride damage. Up to now, only few experiments have been carried out on a restricted number of materials (old bricks [Wendler, 2002] and sandstone containing clay [Snethlage and Wendler, 1997; Wendler and Ruckert-Thumling, 1992]). The research reported in this paper aimed at studying the dilation phenomenon in detail and verifying it for different salt types. The research was carried out in steps. Firstly the effect of NaCl on the hydric and hygric dilation behaviour was investigated. Secondly, to check if the observed behaviour was limited to NaCl or common to more salts, the behaviour of two other salts, NaNO3 and KCl, was studied. These two salts were selected because they are both hygroscopic, they do not have hydrated forms and each of them has one ion in common with NaCl. Finally, the effect on the dilation behaviour of a crystallization inhibitor, modifying the crystallization habits of the salt, was investigated. All experiments were performed on a limecement mortar.

This study was carried out in the EU Research Project COMPASS 'Compatibility of Plasters And renders with Salt loaded Substrates in historic buildings' (EVK-CT 2001-00047).

2 Materials and methods

2.1 Material properties

A lime-cement (lime: cement: siliceous sand = 1:4:20) mortar has been selected for this study. This mortar has been found in previous laboratory experiments to be extremely susceptible to sodium chloride damage, also because of its very low mechanical strength (Table 1). The physical properties of the hardened mortar were determined on 5x5x2cm specimens and are summarized in Table 1. The pore size distribution was determined by Mercury Intrusion Porosimeter (MIP). The mortar has a high porosity mainly constituted by large ($4-7\mu m$) and very large ($80\mu m$) pores. The presence of voids larger than 150, not measured by MIP but visible by Environmental Scanning Electron Microscope (ESEM), may explain the difference in

the open porosity measured by immersion and by MIP respectively.

2.2 Preparation and salt contamination of the specimens

Specimens with a size of 20x10x2cm were used for the study of the hydric and hygric behaviour described in the following sections. Since the physical properties of the mortar are strongly influenced by the suction of the substrate on which it is applied [Wijffels and van Hees, 2000], the plaster slabs were not prepared in a mould but on top of a brick substrate from which they were detached after one day. A filter paper was used between substrate and plaster to facilitate the detachment. The specimens were cured under plastic sheets for 24 hours; then they were detached from the substrate and cured for 15 days at 20°C 65%RH and for the subsequent 15 days at 3% CO₂. After this period the complete carbonation of the mortar was verified by spraying phenolphthalein on a broken section.

Table 1: Physical and mechanical properties of the mortar

Dry density		Porosity		WAC**	Mechanical strength**	
(Kg/m³)		(% by volume)		Kg/(m ² h ^{0.5})	N/mm²	
By immersion*	By MIP	By immersion*	By MIP		Compr.	Flex.
1.795	1.888	32.3	27.6	0.064	0.3	0.1

^{*} Determined according to [van der Klugt, 1994]

Before measuring the hydric and hygric behaviour the specimens were contaminated with NaCl according to the following procedure: the mortar slabs were sealed on the four lateral sides by epoxy resin and a NaCl saturated solution was introduced from the bottom of the specimen (i.e. the side in contact with the filter during preparation) by capillary rise. An amount of solution equal to the Capillary Moisture Content (C.M.C.) of the mortar (i.e. the moisture content enough to wet the upper surface) and leading to 2%m/m of NaCl in the specimen was used. This amount of salt was proved to be enough to damage this material in a previous crystallization test [Lubelli et al, 2006]. After contamination, the specimens were closed at the bottom with removable tape and dried at 10%RH and 20°C until a constant weight was reached. This drying process led to almost no efflorescences; salts accumulated just beneath the evaporation surface of the specimen.

Once the specimens contaminated with salt were dried, metal supports for the Linear Variable Differential Transformer (LVDT) to be used in the measurement of the hydric and hygric behaviour, were glued, at a distance of 150mm from each other, on the surface of the mortar slabs by a two components glue.

^{**} Determined according to [CEN, 1996]

^{***} Determined according to [CEN, 1999]

2.3 Test set-up

The hydric and the hygric behaviour of the specimens were monitored inside a climatic cabinet in which the temperature and RH of the air can be programmed and controlled. In order to reach a very low RH, dry air was blown in the cabinet. The measuring apparatus (Figure 1) was composed by:

- balance with an accuracy of 0.1g, connected to a PC, for monitoring the variation in specimen weight due to hygroscopic moisture uptake and release
- LVDT fixed to the supports glued on the mortar and connected, through the data acquisition system Labview, to a PC
- two data loggers (ESCORT) for double check of the temperature and RH of the air in the climatic cabinet

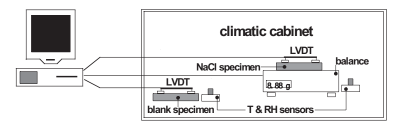


Figure 1: Test set up: the specimens, on which the LVDT are applied, are placed in a climatic cabinet. A PC collects data on the dilation and on the weight changes. T and RH sensors are used for additional check of the environmental conditions

3 Experimental study

3.1 Hydric behaviour of NaCl contaminated specimens

The hydric behaviour was studied both on NaCl contaminated and blank mortar slabs. The specimens were set in the climatic cabinet at 20°C and 50%RH and they were allowed to absorb water by capillary rise from the bottom for a period of two days. Then the bottom of a specimen was sealed with tape, the RH in the cabinet lowered to 0%RH and the drying started. After about 4 weeks, the wet-dry cycle was repeated.

Data on the temperature and RH of the air, as well as the dimensional changes of the specimens at its surface, were collected continuously. Figure 2 shows the results of the experiment. During wetting, both the salt contaminated and the reference specimens have a similar behaviour, consisting of initial shrinkage (due to the wetting of the lower part of the sample) followed by dilation (corresponding to the migration of water to the surface of the sample). During drying,

the blank specimen shrinks, while the salted specimen, after a period of about 4 days in which it does not show any change in size, starts dilating. It can be reasonably supposed that dilation occurs when, due to evaporation of water, the salt starts crystallizing. The maximum dilation is about $0.13\mu\text{m/mm}$ and is reached in 5 days. During the re-wetting both specimens dilate, but the dilation is larger in the salt contaminated than in the blank specimen. During drying the blank specimen shrinks while the salted specimen dilates further.

This experiment shows that the presence of sodium chloride strongly influences the hydric behaviour of this material. The salt contaminated specimen undergoes an irreversible dilation; on the contrary, the displacements of the blank specimen, apart from being smaller, are reversible.

In the described experiment the effect of dissolution and crystallization of salt on the shrinkage/dilation of the material cannot be followed in detail. In fact the amount of absorbed and evaporated water cannot be measured without handling of the specimen; besides it is not known at which moment water, absorbed by capillarity from the bottom of the specimen, reaches the surface where most of the salt accumulates. For a better understanding of the dilation phenomenon, a slower and more controlled moisture absorption and release is necessary. The use of RH changes through the RH of crystallization of NaCl (RH $_{\rm cryst}$ = 75%) allows to reach this aim. In fact hygroscopic moisture adsorption is a slower process than capillary absorption; therefore its effects can be better monitored. In the following section the study of the effect of RH changes on the dilation of salt contaminated specimens is described.

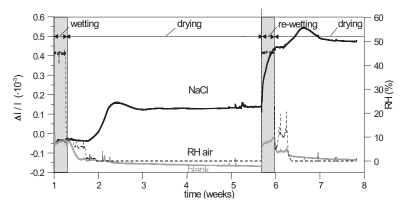


Figure 2: Dilation of blank and NaCl contaminated specimens during wetting at RH=50% (gray areas) and drying at 0%RH (white area) cycles

3.2 Hygric behaviour of NaCl contaminated specimens

The hygric behaviour of NaCl contaminated and blank (reference) mortar slabs was measured. The specimens, prepared as described in section 2.2, were placed in the climatic cabinet. The

temperature was maintained constant at 20°C and the RH was varied between $10\% \pm 2\%$ RH and 92% RH every 48 hours. Six RH cycles were performed. During the test the NaCl contaminated specimen was put on a balance and its weight was continuously monitored. The results, reported in Figure 3, show that the blank specimen dilates during the high RH period and shrinks during the low RH period as expected [Hiller, 1964], while the NaCl contaminated specimen behaves in the opposite way. The dimensional changes of the salt contaminated specimen are not only of opposite sign but also much larger than in the case of the blank specimen. When the cycles are repeated, dilation increases more than shrinkage: this leads to an increasing irreversible dilation that reaches, at the end of the sixth cycle, $1.3\mu\text{m}/\text{mm}$. On the contrary, the dimensional changes are completely reversible in the case of the blank specimen.

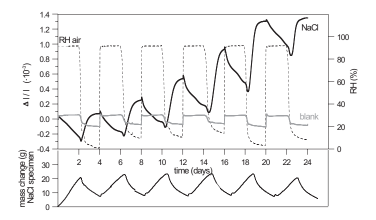


Figure 3: Dilation of blank and NaCl contaminated specimens (upper graph) and weight change of NaCl contaminated specimen (lower graph) during RH cycles

The hygric dilation measured in the NaCl contaminated specimen is considerable, as shown by a comparison with the results obtained in similar experiments performed on other materials and reported in literature (table 2). The dilation leads to a stress that is likely to exceed the low tensile strength of the plaster and in fact results, at the end of the test, in serious damage. The damage consists in sanding of the outer layer of the plaster (Figure 4); the material loss is about 7% of the initial weight of the specimen. The serious damage observed stresses the importance of RH changes for the development of the salt decay in weak lime-cement mortars. Our results are in accordance with the data reported in literature for clay containing sandstone and old brick [Wendler and Ruckert-Thumling, 1992; Snethlage and Wendler,1997; Wendler, 2002]. Moreover, the experiments on the lime-cement mortar allow to definitely assert that the presence of reactive clay is not a necessary factor for the occurrence of irreversible dilation in NaCl contaminated materials.

In order to move a step forward in the understanding of the mechanism, more detailed

experiments were necessary. Performing a RH cycle slowly and with small increments, would allow identifying the point at which dilation and shrinkage occur and relate them unequivocally to either salt crystallization or dissolution. A new experiment was set up in which the RH was varied stepwise between 30 and 96%RH. Each step in RH was done in 1 hour and was followed by 23 hours of equilibration at constant RH. The experimental data (Figure 5) show clearly that the blank specimen dilates while the RH increases and shrinks while the RH decreases. On the other hand the NaCl contaminated specimen shrinks at RH higher than 75% and considerably dilates when the RH drops below the 75%. Note that in the presence of NaCl water condensates when RH > 75%, while in the absence of NaCl it starts to condensate at RH \sim 100%.

Table 2: Hygric dilation in material contaminated with NaCl

Material	Hygric dilation (RH changes)		
Lime/cement mortar without NaCl (this study)	0.13μm/mm (5-90%RH)		
Lime/cement mortar with NaCl (this study)	1.3µm/mm (after 6 cycles 5-90%RH)		
	0.3μm/mm (after 1 cycle 5-90%RH)		
Sandstone with clay without NaCl [Wendler and	0.2μm/mm (35-90%RH) (after 1 or 6cycles)		
Ruckert-Thumling, 1992]			
Sandstone with clay in the presence of NaCl	0.5µm/mm (after 6 cycles 35-90%RH)		
[Wendler and Ruckert-Thumling, 1992]			
Old brick +NaCl [Wendler, 2002]	0.3-4µm/mm (after 9 cycles 40-85%RH)		
New brick + NaCl [Wendler, 2002]	0.1-1µm/mm (after 9 cycles 40-85%RH)		

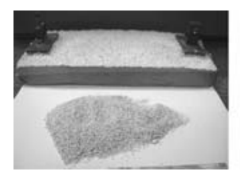




Figure 4: Sanding of the outer layer of the lime-cement mortar in the presence of NaCl after 6 RH cycles

Three parts can be distinguished in the curves:

1. Increasing RH (30-75%RH)

In this range of RH the behaviour of the blank and of the NaCl contaminated specimens is similar: both dilate when the RH increases. The dilation is due to water vapor adsorption [Hiller, 1964].

2. Increasing and decreasing the RH while RH > 75% of NaCl (75-96-75%RH).

After crossing the 75%, the behaviour of the NaCl contaminated specimen starts to deviate from the specimen without salt: as soon as liquid water is present in the plaster (as shown by the weight change) and the salt starts dissolving, shrinkage is measured. The shrinkage of the salt contaminated specimen can be explained by the release of the tensions developed during the previous crystallization of the salt.

3. Decreasing the RH (75%-45%RH)

When the RH drops below the 75%, the salt contaminated specimen starts drying (as shown by the weight loss) and dilates considerably. The curve becomes steeper with decreasing RH. At the end of the test, after 4 days of drying, the dilation measured is about $0.35\mu m/mm$. The not contaminated specimen shrinks, as expected, when the RH is lowered; at the end of the cycle it comes back to the original size.

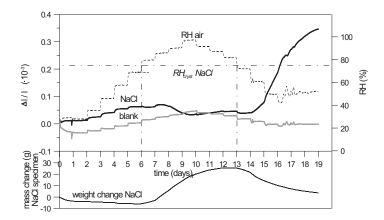


Figure 5: Dilation of blank and NaCl contaminated specimens (upper graph) and weight change of NaCl contaminated specimen (lower graph) during a single RH cycle (max RH 95%)

A further prove that the behaviour of a salt contaminated specimen is modified only if the equilibrium RH of the salt is crossed is given by another experiment in which the RH was varied but kept below 75% (Figure 6): in this case no irreversible displacement took place.

3.3 NaNO3 and KCl

The above-described experiments proved that NaCl is able to produce irreversible dilation during crystallization. To check whether this behavior is specific for NaCl or occurs also in the presence of other salts further experiments were set up. Two hygroscopic salts were selected having respectively the cation or the anion in common with NaCl and no hydrated forms: KCl and NaNO₃.

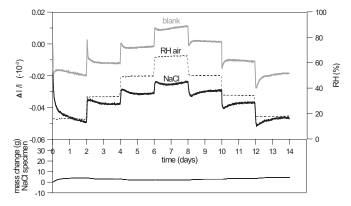


Figure 6: Dilation of blank and NaCl contaminated specimens (upper graph) and weight change of NaCl contaminated specimen (lower graph) during a RH cycle below 75%RH

The specimens were prepared in the same way as for NaCl. The same amount of solution was introduced in the mortar. The concentration of the solution was defined in such a way as to reach in all cases a salt content of 2%m/m of the weight of the dry specimen. As shown in Figures 7 and 8, in both cases shrinkage is observed when the RH exceeds the values where water starts to condensate and the salts dissolve in the water (at 20°C this happens at 85 and 75% for KCl and NaNO₃, respectively [Arnold, 1990]. When the RH decreases below these values the specimens start drying, the salts crystallize and dilation occurs. Therefore it can be concluded that this behaviour is not typical for NaCl but is a more general phenomenon.

3.4 Acid and alkaline pH

The possible effect of acidity or alkalinity of the NaCl solution on the hygric dilation has been investigated by contaminating two specimens with NaCl solutions having a pH of respectively 1.5 and 14. The response of the two specimens to the RH changes is similar and comparable to the one measured on NaCl contaminated specimens: shrinkage is measured during dissolution and expansion during drying (Figure 9). Therefore we conclude that the pH does not play an important role in the studied mechanism. It should also be mentioned that the acid pH is

probably neutralized in the alkaline environment of the mortar.

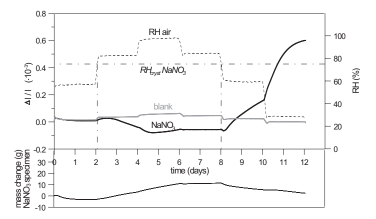


Figure 7: Dilation of blank and NaNO₃ contaminated specimens (upper graph) and change in weight of the NaNO₃ contaminated specimen (lower graph) during a RH cycle

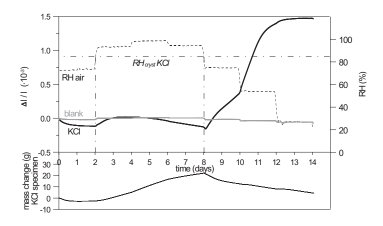


Figure 8: Dilation of blank and KCl contaminated specimens (upper graph) and change in weight of the KCl contaminated specimen (lower graph) during a RH cycle

3.5 NaCl with crystallization inhibitor

The use of crystallization inhibitors may also help answering some questions about the damage mechanisms since they are supposed to increase the supersaturation level (and therefore the theoretical crystallization pressure) at which salt crystallization occurs and modifies the crystal habits [Rodriguez-Navarro, 2002]. Sodium ferro-cyanide ($Na_4Fe(CN)_6$ $10H_2O$) has been chosen since it has proven to be very effective in inhibiting NaCl crystallization [Rodriguez-Navarro et al, 2002].

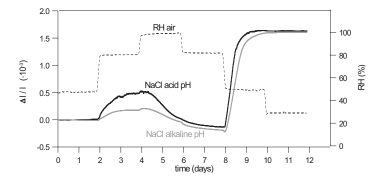


Figure 9: Dilation of specimens contaminated with acid and alkaline NaCl solution during a RH cycle

Two mortar specimens were contaminated with a NaCl saturated solution with 0.1% (m/m) and without sodium ferro-cyanide, $\mathrm{Na_4Fe(CN)_6}$ 10H₂O. In both specimens the same amount of salt was introduced. The salt contamination procedure was the same as for the other specimens, apart for the fact that in this case to prevent efflorescences, which are favored by the presence of the inhibitor, the specimens were dried at 40°C.

The specimens were then placed in the climatic cabinet and the RH was varied step by step between 30 and 98%RH. In Figure 10 the obtained dilations are reported. The specimen contaminated with NaCl solution shows significant dilation during drying, whereas the specimen with NaCl plus inhibitor displays almost no change in size. The almost complete drying out (and therefore the necessary occurrence of salt crystallization) of the specimen containing the inhibitor can be checked by looking at its weight at the end of the experiment.

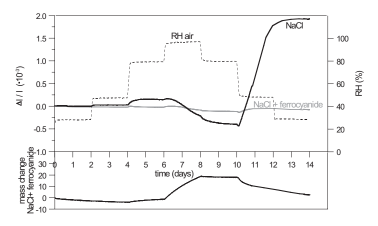


Figure 10: Dilation of NaCl contaminated specimen with and without inhibitor (upper graph) and weight change of NaCl contaminated specimen with ferrocyanide (lower graph) during a RH cycle

In order to investigate the long-term effect and the consequences of the presence of the inhibitor on the decay, another experiment, consisting of repeated dissolution/crystallization cycles, was performed. Also in this case NaCl contaminated specimens, with and without inhibitor, were used. The air RH varied between 0 and 93%. The results obtained in this experiment (Figure 11) in the presence of ferrocyanide confirm the previous ones: the specimen dilates during dissolution and shrinks during crystallization of the salt and its dimensional changes are reversible. The specimen contaminated with NaCl behaves as expected (shrinks during dissolution and dilates during crystallization).

At the end of the 5th cycle the NaCl contaminated specimen show little sanding of the outer surface, whereas the specimen containing the inhibitor did not suffer any damage. The absence of damage, in spite of the high degree of supersaturation and therefore crystallization pressure at which the salt is expected to crystallize, cannot be explained (like in [Rodriguez-Navarro et al, 2002]) by the increased tendency of the salt to efflorescence. In fact at the end of the experiment only few salts were visible on the surface.

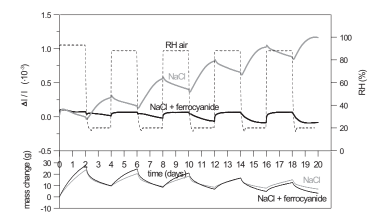


Figure 11: Dilation (upper graph) and weight change (lower graph) of NaCl contaminated specimen with and without inhibitor during five RH cycles

4 ESEM observations

Environmental Scanning Electron Microscopy (ESEM) studies have been performed on the specimens subjected to RH cycles, after their complete drying.

The cross sections of the outer layer (about 3mm thick) of mortars contaminated with NaCl, $NaNO_3$, KCl and NaCl plus inhibitor were observed using a Back Scattered Electron (BSE) detector. The composition of the salt crystals was checked with Energy Dispersive Spectroscopy X-ray Microanalysis (EDX). The investigations aimed to study the location and the habits of salt crystallization in pores.

In the ESEM picture of the lime-cement mortar contaminated with NaCl, the salt is visible in larger pores, creating a layer over the pore walls (Figure 12). It looks as if a strong interaction exists between the material and the salt crystals. This is in accordance with the observations reported in other studies [Benavente et al, 2004]. Under the salt layer, most of the distinguishable pores appear to be empty (Figure 13).

The presence of acid or alkaline pH does not affect the crystallization habit of sodium chloride (Figure 14) in the studied lime-cement mortar.

Crystallization habits similar to the one observed for NaCl were found also in case of $NaNO_3$ and KCl (Figure 15).

In case the crystallization inhibitor was added to the salt solution a different crystallization habit of the NaCl was observed. The salt does not show anymore a strong affinity with the substrate: it does not cover the pore walls in the form of a layer, but crystallizes mainly as agglomerations of crystals, not strongly attached to the material. The salt is not homogeneously distributed in the contaminated area but accumulates in definite zones (Figure 16).

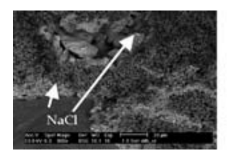
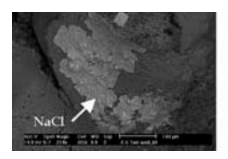




Figure 12: ESEM photomicrographs showing NaCl (lighter areas) crystallizing as a layer over the pore walls of the binder

Figure 13: ESEM photomicrograph showing that, under the salt layer (white area), the pores of the material are not filled with salt



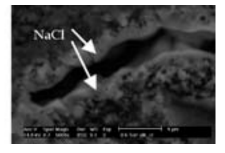
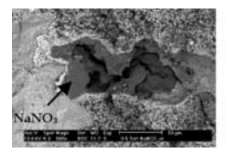


Figure 14: ESEM photomicrographs showing NaCl (lighter areas) crystallizing as a layer in specimens contaminated with acid (left) and alkaline (right) NaCl solution



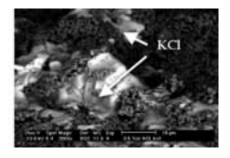


Figure 15: ESEM photomicrographs showing $NaNO_3$ (left) and KCl (right) crystallizing as a layer on the pore wall

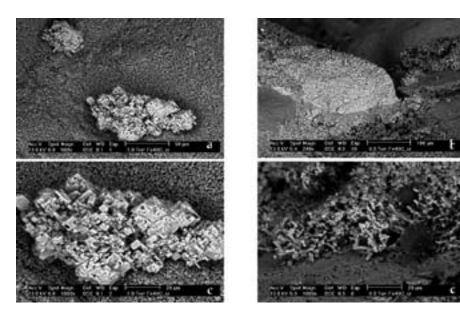


Figure 16: ESEM photomicrograph showing NaCl crystallization in presence of inhibitor; crystal agglomerations (white areas) are observed (a-b); the crystal may be more (c) or less regular (d)

5 Discussion and conclusions

Our experiments unambiguously proof that NaCl is able to modify the hydric and hygric behaviour and the related dilation of a material completely, even in the absence of reactive clay. In the lime-cement mortar NaCl produces dilation during the drying phase of the RH cycle, when the salt crystallizes. The dilation observed in the NaCl contaminated specimens is irreversible and increases with repeated RH cycles until damage occurs. Damage appears as sanding of the outer layer of the specimen where most of the salt has accumulated. It has been proven that this dilation is due to crystallization since it only occurs in presence of salts, when

the RH of the air is low enough to drive the evaporation of water.

The same effect of salt has also been observed for other hygroscopic salts (NaNO₃ and KCl), showing that this phenomenon is not restricted to NaCl but is more general.

The presence of a crystallization inhibitor also modifies the hygric behaviour of NaCl contaminated specimens: no more relevant dilation is observed during drying and crystallization.

The ESEM investigations have shown that NaCl, as well as $NaNO_3$ and KCl, crystallize as a layer on the pore walls. When a crystallization inhibitor is added to the NaCl solution, the salt crystallizes as crystals not adhering to the pore walls.

Combining the ESEM observations with the dilation occurring during crystallization, leads to a theory in which the damage mechanism is based on a mechanical interaction between salt and pore wall. Salts crystallizing as a layer on the pore walls seem to be able to strongly affect the hygric dilation of the material producing relevant expansion during drying of the specimen and crystallization of the salt. It seems that these salt layers are able to transfer stress to the pore walls and thereby causing dilation (Snethlage and Wendler 1997; Puhringer et al 1985). A similar effect would then not be possible when the same salt crystallizes without adhering to the material, as in the case in which the crystallization inhibitor is added. Further experiments have been set up in order to check this hypothesis.

Acknowledgments

The authors wish to thank the technician Gerrit van der Ende for the set up of the measuring apparatus. The research was developed in the framework of the EU project COMPASS, (Compatibility of Plasters and Renders with salt loaded substrates in historic buildings), EU Contract no. EVK4-CT-2001-0047-DGXII www.compass-salt.org.

References

- Arnold A. (1990) 'Salt weathering on monuments' in F. Zezza (Ed.), Advanced Workshop Analytical methodologies for the investigation of damaged stones, pp.31-58
- Benavente D., Garcia del Cura M.A., Garcia-Guinea J., Sanchez-Moral S. and Ordonez S. (2002) 'Role of pore structure in salt crystallization in unsaturated porous stone', in *Journal of Crystal Growth*, Nr.260, pp. 532-544
- Correns C.W. (1949) 'Growth and dissolution of crystal under linear pressure', in *Discussion of the Faraday Society*, Vol. 5, pp.267-271
- European Committee for Standardization (CEN) (1996) European Standard prEN 1015-18 Methods of test for mortar for masonry. Determination of water absorption coefficient due to capillary action of hardened rendering mortar

- European Committee for Standardization (CEN), (1999) European standard EN1015-11, Methods of test for mortar for masonry Part 11: Determination of flexural and compressive strength of hardened mortar
- Flatt R.J. (2002) 'Salt damage in porous materials: how high crystallization pressure are generated' in Journal of Crystal Growth, Nr.242, pp. 435-454
- Hiller K.H. (1964) 'Strength Reduction and Length Changes in Porous Glass Caused by Water Vapor Adsorption' in *Journal of Applied Physics*, Vol.35, pp. 1622-1628
- van der Klugt L.J.A.R. and Koek J.A.G. (1994) 'De kwaliteit van voegen in metselwerk' in SBR-publication, Nr. 299, pp. 80-81
- Lubelli B., van Hees R.P.J. and Groot C.J.W.P. (2006) 'The effect of environmental conditions on the decay due to sodium chloride: a step in the development of an effective weathering test' in *Studies in Conservation*, 51 (1)
- Pel L., Huinink H.P. and K. Kopinga (2002) 'Ion transport and crystallization in inorganic building materials as studied by nuclear magnetic resonance' in *Applied Physics Letters*, 81, pp. 2893-2895
- Puhringer J., Berntsson L. and Hedberg B. (1985) 'Hydrate salts and degradation of materials' in G. Felix (Ed.), *Proceedings of the 5th International Congress on Deterioration and Conservation of Stone*, pp. 279-288
- Rijniers L.A, Huinink, H.P., Pel L. and Kopinga K., (2005) 'Experimental Evidence of Crystallization Pressure inside Porous Media' in *Physical Review Letters*, 075503 94
- Rodriguez Navarro C., Linares-Fernandez L., Doehne E. and Sebastian E. (2002) 'Effects of ferrocyanide ions on NaCl crystallization in porous stone' in *Journal of Crystal Growth*, Issue 3-4, Nr.243, pp.503-516
- Scherer G. (1999) 'Crystallisation in pores' in Cement and Concrete Research, Vol.8, Nr.29, pp. 1347-1358
- Snethlage R. and Wendler E. (1997) 'Moisture Cycles and sandstone degradation' in Baer N.S. and Snethlage R. (Eds.) Saving our architectural heritage: the conservation of historic stone structures, pp. 7-24
- Wendler E. and Ruckert-Thumling R. (1992) 'Gefuezerstorendes Verformungsverhalten bei Salzbefrachteten Sandsteinen unter hygrischer Wechselbeanspruchung' in Wittman F.H. (Ed.) Werkstoffwisseschaften und Bausanierung, Vol.3, pp. 1818-1830
- Wendler E. (2002) 'Laboratory measurements on salt-loaded brick samples in periodically changing climate conditions' in von Konow T. (Ed.) *The study of salt deterioration mechanisms*, pp.81-87
- Wijffels T.J. and van Hees R.P.J. (2000) 'The influence of the loss of water of the fresh mortar to the substrate on the hygric characteristics of the so-called restoration renderings', in Verhoef L.G.W. and Wittmann F.H. (Eds.) *Maintenance and re-strengthening of materials and structures Plaster*, pp. 49-54