Pore Structure and Water Transfer in Pietra d’Aspra Limestone: A Neutronographic Study

Luciana Randazzo 1, Giuseppe Paladini 2, Valentina Venuti 2,*, Vincenza Crupi 3, Frédéric Ott 4, Giuseppe Montana 5, Michela Ricca 1, Natalia Rovella 1, Mauro Francesco La Russa 1,*, and Domenico Majolino 2

1 Department of Biology, Ecology and Earth Science, University of Calabria, via Pietro Bucci cubo 12B piano 2, 87036 Arcavacata di Rende (CS), Italy; luciana.randazzo@unical.it (L.R.); michela.ricca@unical.it (M.R.); natalia.rovella@unical.it (N.R.)
2 Department of Mathematical and Computer Sciences, Physical Sciences and Earth Sciences, University of Messina, Viale Ferdinando Stagno D’Alcontres 31, 98166 Messina, Italy; gpaladini@unime.it (G.P.); dmajolino@unime.it (D.M.)
3 Department of Chemical, Biological, Pharmaceutical and Environmental Sciences, University of Messina, Viale Ferdinando Stagno D’Alcontres 31, 98166 Messina, Italy; vcrupi@unime.it
4 Laboratoire Léon Brillouin, CEA Saclay, 91191 Gif-sur-Yvette CEDEX, France; frederic.ott@cea.fr
5 Department of Earth and Sea Sciences (DiSTeM), University of Palermo, Via Archirafi 22, 90123 Palermo, Italy; giuseppe.montana@unipa.it

* Correspondence: vvenuti@unime.it (V.V.); mlarussa@unical.it (M.F.L.R.)

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Abstract: Neutron radiography (NR) was here applied to study the effects of two different commercially available consolidants on the water absorption properties in a particular type of limestone (biocalcarenite), known as Pietra d’Aspra stone, which is one of the most extensively used lithotypes in Sicilian Baroque buildings. Our attention was mainly focused on the evaluation, using a fast and nondestructive visualization of water motion through capillarity, of the effectiveness of such layers as consolidating agents in view of preserving and maintaining both old and modern structures. The biocalcarenite was treated with nanosilica (Nano Estel®) and nanolime (CaLoSil®) by brushing it until full saturation, and then artificially weathered by salt crystallization and temperature/relative humidity jumps. Liquid distribution and height of the water front were monitored as a function of time. Significant differences in the water suction behavior among all the investigated samples were highlighted, which allowed us to gain insight into the coating/substrate interaction mechanisms which regulate the fluid mobility inside the porous network of the limestone.

Keywords: neutron radiography; porous limestones; consolidants; water kinetics; artificial weathering; capillarity

1. Introduction

The presence of water inside the porous matrix of building materials, whether artificial (concretes and mortars) or natural (stones), plays a main role in determining their mechanical performance and stability. As well known, salt crystallization and transportation of unwanted substances through water penetration can lead to the formation of cracks and plane distortions, strongly affecting the structural integrity of the construction itself. Restoration treatments aimed at “conservation” should preserve the original material while slowing down the decay process as much as possible. Stone durability is a term describing how well the material retains its original physical and mechanical properties. As a consequence, it concerns several aspects such as texture, structure, and mineralogical composition, as well as methods of quarrying and handling, applied loads, and environmental conditions to which
the considered stone has been exposed [1–6]. In particular, during the past few centuries, porous calcareous stones were extensively used as building material in several areas of the Mediterranean basin. In coastal urban environments, after a protracted period of exposure to the atmospheric agents, these lithotypes are very often subjected to extensive crumbling due to soluble salt crystallization in pore spaces [7–13]. Consequently, for this type of natural building material, the relationship among pore structure, water/moisture transport, and salt crystallization has been extensively investigated for understanding the mechanisms and kinetics of degradation processes [2,9,14–19].

In this framework, different consolidant products have been commercialized and used in the last few years in the treatment of building materials of historical monuments, in order to preserve the whole structure and to guarantee a longer durability [20]. The high efficiency and minimum appearance alterations of such consolidating agents represent two highly demanded conditions in the field of conservation science. Both aspects depend on how these products interact with the relative substrates, affecting the penetration depth and the difference in porosity between the consolidant and the underlying bulk stone. In particular, the 3D pore distribution of both the stone and the coating layer represents a crucial parameter since it regulates the fluid mobility inside the material.

Currently, liquid imbibition dynamics can be visualized through different approaches including gamma rays [21,22], nuclear magnetic resonance (NMR) [23], and neutron radiography (NR) [24]. However, it must be pointed out that NR, among the other techniques, represents the best solution for the evaluation of the height of the water front and its 3D distribution, as all hydrogen-rich fluids are characterized by higher neutron attenuation coefficients with respect to elements typically present in limestone specimens. The study of water migration by capillarity in calcite-based materials (such as limestones) by means of NR has gained considerable attention by the scientific community in recent years [25–30]. It has been successfully applied for the study of water movement by spontaneous imbibition in rocks [31–33], sandstones [34,35], concrete [36,37], and bricks [38]. Generally, a linear trend of the height of the water front (h(t)) with the square root of the exposure time is attained, according to the Lucas-Washburn (L-W) equation. It is worth noting that the water rise by capillarity follows the aforementioned classical behavior only for structures characterized by a homogeneously distributed porosity. However, violations in the diffusion-like process can be often achieved [39,40], at the early stage of water contact or after a certain period of time, revealing the occurrence of several processes during the water suction and/or the presence of nonuniform/disordered porosimetry within the inner structure of the investigated sample.

In the present work, neutron radiography was successfully applied in order to quantify, in a nondestructive way, the susceptibility to weathering of a particular type of limestone, i.e., Pietra d’Aspra stone. This variety of biocalcarenite was extensively used in the architectural practice of Palermo (Sicily, Italy) and its surroundings, since antiquity and especially in the second half of the 17th century, for building Baroque palaces, nobiliary villas, and churches [41]. The generalized interest in this type of porous calcareous stone derives from its availability in the territory and from the relative ease of quarrying and workability. Its main forms of deterioration are represented by blackish sulfur-bearing encrustations developed on surfaces protected by rain-washing, or alveolar decohesion and extensive crumbling in the exposed parts of the masonry which are related to the precipitation of soluble salts [42]. On the basis of the intrinsic value and the number of monumental complexes made with Pietra d’Aspra stone, the development of innovative experimental procedures aimed at evaluating performances of consolidating treatments on this specific limestone is considered, at present, of great attraction and relevance.

More specifically, in our case, the interaction mechanisms between two consolidant products, namely, nanosilica (Nano Estel®), nano-SiO₂, NS) and nanolime (CaLoSil®), Ca(OH)₂, NL), and the analyzed stone were mainly addressed, with the aim of highlighting the water transport behavior exhibited by the investigated material after consolidation. Treated samples were also exposed to different accelerated weathering tests such as 15 cycles of salt crystallization and temperature/relative humidity jumps, in order to simulate the material’s response to different natural stresses. Results
reported here could be used, in principle, to propose an optimized procedure for preserving and maintaining buildings/objects of interest in the field of cultural heritage and conservation science.

2. Materials and Methods

2.1. Geological Notes and Sampling

Biocalcarenite outcrops have been intensively quarried in the whole plain of Palermo throughout past centuries [43–45]. These deposits can be regarded as deposits of shallow tidal sea/representing low sea-level sediments over the Tyrrenhian period (Upper Pleistocene, 0.5–0.1 Ma), and they overlie the Mesozoic Carbonatic Formation and/or Numidian Flysch Formation with discordance [46]. This lithotype, locally called Pietra d’Aspra, comes from quarrying districts located a few kilometers west of Palermo. Ancient quarries can still be observed. Different varieties belong to the same calcarenite lithotype, characterized by a different compaction following the deposition, a different carbonate cement percentage, and, consequently, having a different total open porosity, grain distribution, and color. With regard to these parameters, it is possible to distinguish lithotypes with a grain size ranging from medium to fine arenite (0.5–0.125 mm) and lithotypes with a grain size ranging from coarse to medium arenite (2–0.5 mm), more or less cemented. In particular, the coarse-grained calcarenite, well cemented and characterized by a high value of compressive strength (30 MPa), can mostly be used for walls and facades such as basements and columns. On the other hand, soft calcarenites were used as decorative materials thanks to their easy workability and low value of compressive strength [45,47].

Representative samples of a coarse-grained calcarenite were chosen for this study. Calcarenite samples were taken from the Guttadauro quarry front located in Via Cotogni, Aspra village, Palermo (Figure 1a,b).

\[
\gamma = \frac{2\gamma \cos \theta}{P}
\]

\(r=\frac{2\gamma \cos \theta}{P}\), where \(r\) is the pore entry radius in which mercury is intruded, \(\gamma\) is the surface tension, and \(\theta\) and \(P\) represent the contact angle of mercury with the solid and the applied pressure, respectively. After this preliminary

\(\theta\)

\(P\)

\(\gamma\)

\(r\)

\(\theta\)

\(P\)
characterization, samples (dimensions $\sim 5 \times 5 \times 2$ cm$^3$) were treated with the two commercial products, i.e., Nano Estel® (CTS srl, Vicenza, Italy) and CaLoSiL® (Bresciani srl, Milano, Italy), whose applications were carried out on the surface of each sample by brushing it several times until saturation. After a curing time period of 28 days, accelerated weathering tests (15 cycles of salt crystallization and temperature/relative humidity (T/RH) jumps) were carried out on treated and untreated samples (see Table 1), following the procedure reported in [48]. In particular, each cycle of salt crystallization consists of (a) 2 h of immersion in a supersaturated solution of sodium sulfate (14% w/w at 20 °C) for 10% of the height, (b) 8 h of drying in an oven at 45 °C, and (c) 16 h of cooling at room temperature. Initial and final weights after each cycle were measured for each test sample (in triplicate). The resulting weight loss (average value of three measurements) was, therefore, determined. Lastly, three specimens for each treatment and three specimens for untreated ones underwent a humidity/temperature jump in a climatic chamber after three cycles of salt crystallization with the following conditions: temperature (T) variation between 20 °C and 45 °C and relative humidity (RH) variation between 40% and 80% for about 1500 h.

| Sample Code | Products       | Experimentation Tests | Preliminary Characterization (PLM, MIP) | Weight Loss Measurements | Neutron Radiography Analysis |
|-------------|----------------|-----------------------|----------------------------------------|-------------------------|----------------------------|
| PA1         | Nano Estel®    | Climatic chamber (T/RH) after 3 cycles of salt crystallization | -                                       | $\times (3)$             | $\times (1)$               |
| PA2         | Nano Estel®    | Salt weathering (15 cycles) | -                                       | $\times (3)$             | $\times (1)$               |
| PA3         | Nano Estel®    | Not weathered         | -                                       | -                       | $\times (1)$               |
| PA4         | CaLoSiL®      | Climatic chamber (T/RH) after 3 cycles of salt crystallization | -                                       | $\times (3)$             | $\times (1)$               |
| PA5         | CaLoSiL®      | Salt weathering (15 cycles) | -                                       | $\times (3)$             | $\times (1)$               |
| PA6         | CaLoSiL®      | Not weathered         | -                                       | -                       | $\times (1)$               |
| PA-TQ       | Untreated      | Not weathered         | $\times (1)$                           | -                       | $\times (1)$               |
| PA-TQ1      | Untreated      | Climatic chamber (T/RH) after 3 cycles of salt crystallization | -                                       | $\times (3)$             | -                          |
| PA-TQ2      | Untreated      | Salt weathering (15 cycles) | -                                       | $\times (3)$             | -                          |

After salt weathering and T/RH jumps, a preliminary evaluation of the consolidating action of such coatings was achieved by comparing the weight loss in treated and untreated specimens through standard laboratory-based methodologies.

2.3. Neutron Radiography Analysis

One test specimen for seven samples (see Table 1) was used for neutron radiography analysis. Neutron radiographs were collected at the cold neutron imaging spectrometer IMAGINE of the Laboratoire Léon Brillouin (LLB) at the Orphée Reactor, in Saclay (F). By increasing the $L/D$ ratio (with $L$ equal to the distance between the entrance aperture of the neutron beam and the image plane (400 cm) and $D$ equal to the diameter of the collimator aperture (1 cm)) up to 400, absorption contrast values with high resolution ($\sim 0.1\%$) were achieved, providing a spatial resolution of about 250 µm. The incoming beam, characterized by a wide spectrum of neutrons having wavelength between 3 and 20 Å, was of $\sim 2 \times 10^7$ neutrons·cm$^{-2}$·s$^{-1}$. Finally, for our measurements, the sample–detector distance was 10 cm.

A sCMOS (scientific Complementary Metal–Oxide–Semiconductor) ANDOR NEO camera coupled to a Canon EFS 60mm F/2.8 Macro USM (Ultrasonic Motor) was used to collect the transmitted beam coming from the sample. The key elemental composition of the investigated limestones, mostly characterized by C, O, Ca, etc., whose neutron cross-sections are typically two order of magnitudes lower than that of hydrogens (H) in liquid media, allows easily following the water motion by
spontaneous imbibition. As a result, even for a low quantity of penetrated water, a good contrast between the solid and liquid phase can be observed.

The obtained raw images were corrected using dark-field \(I_{(df)}\) and open-beam \(I_{(ob)}\) images taken prior to the sample measurements.

Investigated limestones were contained one by one within an aluminum holder and placed onto a 10 mm thickness filter paper pack (Kaltek S.r.l. Padova Italy), in order to scan the entire volume. The dynamics of water suction by capillarity was recorded by stepwise manual addition of water into the aluminum basement, which ensured the saturation of the filter paper pack (Scheme 1).

For each sample, images were recorded with an exposure time of 40 s, until full saturation, except for PA4, for which an exposure time of 10 s was used. In those cases, in which the saturated condition was not completely reached, measurements were collected for no longer than ~20 h.

A homemade macro of the IMAGINE beamline was used to pre- and post-process raw images. Accordingly, neutron radiographs of the dry \(I_{(d)}\) and wet \(I_{(w)}\) stone were corrected to both \(I_{(df)}\) and \(I_{(ob)}\) using the following relationships:

\[
I_{w,\text{corr}} = C \frac{I_{(w)} - I_{(df)}}{I_{(ob)} - I_{(df)}},
\]

\[
I_{d,\text{corr}} = C \frac{I_{(d)} - I_{(df)}}{I_{(ob)} - I_{(df)}},
\]

where \(C\) is a rescaling factor accounting for the neutron beam fluctuations. The removal of unwanted bright pixels coming from scattered \(\gamma\)-rays was achieved by encountering the obtained images through a median filter. The procedure adopted to follow the water content from transmission radiographs consisted of normalizing wet images with respect to dry ones taken prior to the manual addition of water into the aluminum basement. In this way, the contribution of the stone is removed, and the obtained transmitted intensity can be directly related to the amount of water within the stone \([32]\). However, in order to get quantitative information on the amount of water contained within the material, a calibration of the collected neutron transmission values was necessary. Accordingly, a staircase-like sample holder, containing 12 different water contents with standardized thickness (ranging from 0.09 mm up to 5.00 mm), was used to evaluate the neutron transmission vs. water thickness plot. The obtained calibration curve (see Figure 2) follows a logarithmic behavior in the whole range of standardized water thickness used. From the fitting procedure, the value of the water attenuation
Experimental value

\[ I(x, y) = I_0(x, y) e^{-\mu_s \theta_s - \mu_w \theta_w} \]

where \( I_0(x, y) \) represents the intensity of the incoming neutron beam, \( \mu_s \) and \( \mu_w \) are the stone and water attenuation coefficients, and \( \theta_s \) and \( \theta_w \) are the thickness of the stone and the equivalent water thickness.

Since the thickness of the stone is known, the mean moisture content (MC) was firstly evaluated as

\[ MC = \frac{\theta_w \cdot \rho_w}{\theta_s}, \]  

where \( \rho_w \) represents the density of water. Then, by dividing the MC by the density of the stone \( \rho_s \), the water content (WC), expressed as a weight percentage (%), could be easily calculated as

\[ WC = \frac{MC}{\rho_s}. \]

As the investigated stone was mostly composed of calcite (90% of CaCO\(_3\)), \( \rho_s \) was taken to be almost equal to 2.69 g/cm\(^3\).
3. Results and Discussion

3.1. Petrographic and Physicochemical Analysis

*Pietra d’Aspra* can be classified as a grainstone [51] or biosparite [52]. The grain size distribution is quite heterogeneous (Figure 3a) and it is composed of at least two secondary modes, one of them being coarse-grained sand. The principal components are calcareous bioclasts, including fragments of molluscan shells, bryozoans, echinoderms, corals, and foraminifera sized up to 1-1.5 mm (Figure 3b).

![Figure 3. (a) Fresh cut surface of a representative specimen of Pietra d’Aspra (not consolidated and not weathered, sample PA-TQ); (b) thin-section microphotograph of the same sample (scale bar = 0.5 mm).](image)

The most abundant accessory components are lithic fragments (0.1 mm in size), possibly derived from Mesozoic limestones and dolostones, and monocrystalline quartz grains ranging from 0.04 to 0.2 mm in size. More sporadically, clasts of siliceous rocks (quartzarenites and radiolarites) may occur. The micrite syngenetic matrix is, in general, poorly abundant, while different types of diagenetic carbonate cements have been reported, such as “meniscus” and “dog teeth”. Predominantly intergranular macroporosity with cavities having a variable and irregular shape, ranging in size from 0.2 to 5 mm, has also been highlighted. Table 2 summarizes most petrographic and mineralogical characteristics of this stone together with the main physical parameters [53].

| Classification (Dunham/Folk) | Mean Grain Size (mm) | Colour (Munsell) | *CaCO3 | *Quartz | *Dolomite | *HCl-Insoluble Residue (W%) | *ρs |
|-----------------------------|----------------------|-----------------|--------|---------|-----------|----------------------------|------|
| BS/G                        | 2.4                  | 10YR 7/6        | 97%    | 2%      | 1%        | 5                          | 2.69 |
| *SiO2                        | 2.16%                | 0.03%           | 0.55%  | 0.52%   | 0.09%     | 0.78%                      | 52.54% |
| *Na2O                        | < 0.01%              | 0.05%           | 0.05%  | 312 ppm | 2 ppm     | 15 ppm                     | 43.23% |

Legend: BS = biosparite; G = grainstone; ρs = density of the stone (g/cm³). * Data sourced from [53].

As well known, mercury intrusion porosimetry does not allow determining the total volume of the stone; nevertheless, porosimetric curves give useful information about pore size distribution in the range of 200 μm–0.1 μm. The evaluation of the pore range, influencing the water uptake of the calcarenite, is an important factor to understand water transport properties.

Grain size, grain packing, and grain contact usually control pore shape and pore size and, therefore, determine the physical properties related to water transport and storage. Pore shape seems to be of primary importance in the relationship between dynamic and static moduli of elasticity and strength.
It appears that textural properties may represent the key parameters controlling physicomechanical properties and weathering behavior.

The porosimetric curve obtained for the PA-TQ sample (not consolidated and not weathered) reveals that the investigated calcarenite has an important porosity and a very wide range of pore sizes (Figure 4), highlighting the heterogeneity of pore network systems in these types of rocks. This characteristic depends on the differences in grain size and cementation degree. The pore network system is addressed in the migration of water solutions, even if the role of each dimensional range in the capillary transfer is dependent on the pore’s size, shape, and connectivity. The sample is characterized by a bimodal pore network system in which the transfer of water solutions occurs heterogeneously.

Degradation forms, such as alveolization and differential erosion, develop more easily in the varieties of rocks with such a heterogeneous porous network. This heterogeneity makes difficult the migration of the saline solutions through the porous capillary network, mainly because of the poor connection between the pores. A slowdown in the transfer kinetics of the aqueous solutions toward the external surface occurs, determining a retreat of the evaporation front toward the inside of the rock. In these areas, the salts can crystallize and concentrate, causing the granular disaggregation of the rock structure.

After the salt crystallization test, the weight variations of consolidated (PA2 and PA5) and untreated specimens (PA-TQ2) were measured (Figure 5a). A great number of efflorescences, due to the transfer of the salt solution, are more evident on the surface of untreated sample with respect to the consolidated ones. In fact, the presence of the consolidants seems to affect the evaporation rate of the solution at the surface of the stone. The development of efflorescence and sub-efflorescence leads to an increase in weight of the individual samples, which seems to balance the slight loss of material suffered by the specimens. This behavior is not dependent on the type of used consolidant, although minor variations occurred in the specimens treated with NS. After 15 cycles, a slight smoothing of the edges was highlighted in all the specimens, especially in the unconsolidated ones, followed by the samples treated with the NL product. For what concerns consolidated specimens subjected to humidity changes along with temperature fluctuations (PA1 and PA4), the weight variations were also recorded during the first three cycles of salt crystallization and at the end of the stay inside the climatic chamber.
(~60 days), and they were finally compared with those of untreated samples (PA-TQ1). Figure 5b shows that the greatest weight gain occurs for the samples treated with NL, indicating that specimens became heavier from salt precipitation compared to both the untreated and the NS-treated ones. However, none of the rock samples showed obvious cracks, flaking, or loss of material, even at the end of the experiment, meaning that the damage induced by subsequent changes in temperature and humidity seems to be rather weak.

Figure 5. Weight variation (%) (a) after salt crystallization test and (b) after temperature/relative humidity jumps.

3.2. Neutron Radiography Analysis

Time-lapsed radiographs of water capillary suction collected for PA1 sample at nine different selected time-steps are displayed in Figure 6, as an example.

Figure 6. False-colored transmission images of water imbibition in PA1 sample at (a) 40 s, (b) 2560 s, (c) 3840 s, (d) 5120 s, (e) 6400 s, (f) 7680 s, (g) 8960 s, (h) 10,240 s, and (i) 11,560 s. In green, the 5 × 5 cm² surface area of the limestone is displayed, whereas the blue/purple/red regions account for the liquid phase characterized by an ever-increasing amount of water. The quantitative evaluation of the water content is addressed from the analysis of the wetting profiles (see text for details).
A first inspection of Figure 6 reveals how the water rising in PA1, as for the majority of the investigated samples, is characterized by an extremely fast “rate” of spontaneous imbibition during the first stage of the absorption test, followed by a slow dynamic. This evidence, together with a nonuniform distribution of water during the uptake process, supports the well-known heterogeneity and poor interpore connectivity of the analyzed stone, as also revealed by the pore size distribution analysis (see Figure 4). This result can be considered, in a way, experimental evidence of the effect that the bimodal distribution of porosity in Pietra d’Aspra limestone exerts on the water transfer behavior. In fact, as the stone is not uniform whether in terms of porosity or in terms of the size of the individual natural components (i.e., calcareous bioclasts and lithic fragments), a regular rise, as observed in a large variety of similar systems [54], was not expected. Furthermore, as the water kinetics shown in Figure 6 is descriptive of the PA1 sample, consolidated and subjected to a climatic chamber, a difficulty in the migration of saline solutions through the capillary porous lattice can be hypothesized. As a result, salts may shrink and crystallize, leading to a partial granular stone disaggregation. However, as we see later in the text, the presence of nano-SiO$_2$ brushed onto the limestone surface strongly reduces the amount of water (wt.%) within the stone network with respect to that observed for the reference sample (PA-TQ), proving a remarkable effectiveness of such a coating as a consolidating agent against accelerated weathering tests. Moreover, after a time interval of ~2600 s, a slight formation of salt efflorescence can be detected as a blue spot that stretches out over the lateral surface of the stone in the bottom-right area of each image (except for Figure 6a). This evidence reveals the development of a preferential pathway for the liquid water during the imbibition toward the right-lateral surface of the stone, which spilled out after about 2600 s.

In order to provide a quantitative picture of the effects that consolidants exert on the behavior of the water suction through capillarity in our investigated materials, in Figure 7, typical plots of water content (WC) as a function of the sample height are displayed.

A first inspection of Figure 7 reveals that, for all the investigated samples, enhanced water penetration during the first stage of water contact is achieved, as suggested by the diminishing of the distances between two consecutive curves as time increases. This phenomenon can be associated with the presence of diagenetic cement, which slows the absorption process, in conjunction with a progressively diminishing of the “free” space available for water upon approaching the top of the material.

Concerning the reference sample (PA-TQ), the obtained values of water content and height were found to be equal to ~3 wt.% and ~50 mm, respectively. These values are always higher than or comparable to (except for the water content in PA4) those obtained for all the treated and artificially weathered stones, revealing (i) experimental evidence of the interaction between the coating and the substrate, and (ii) the presence of fractures and distortions as a result of microclimatic stimuli and salt crystallization. This occurrence, however, can be considered as proof of the consolidating action of both used products. More in detail, as evidenced by the height reached by water during the first 13,310 s of suction, samples treated with nano-SiO$_2$ (NS) show higher performances with respect to those treated with nanolime (NL). In fact, whereas, in samples PA4, PA5, and PA6 (NL-treated), values of water height and content similar to those observed for the reference sample (PA-TQ) were obtained, in PA1, PA2, and PA3 (NS-treated), the water height reached ~20, ~15 and ~30 mm, respectively, showing a hindering in the water uptake of ~60%. In this case, it is reasonable to assume that the presence of a millimeter-thick layer of NS brushed onto the limestone surface leads to an extensive occlusion of the outer pores, preventing the penetration of water inside the stone network. On the other hand, we can state that the presence of NL, as also occurs in the reference (PA-TQ) specimen, allows the pore network to “breathe”, resulting in a faster water suction kinetics. It is worth remarking that if, on one hand, the almost total obstruction of pores represents a suitable condition in order to avoid the introduction of external unwanted substances through liquid inclusion, on the other hand, an almost open structure, as that resulting from the application of NL, may reduce the risk of formation of cracks and fractures originating from the development of high crystallization pressures.
Water content profiles vs. sample height as obtained following the procedure described in Section 2.3. As already stated, neutron radiographs were collected every 40 s for all samples except for PA4, for which a time-step of 10 s was used. In our case, sample PA4 was measured for a total of 13,310 s, during which a water-saturated condition was achieved. For all the other samples, longer exposure times were necessary, since a slower water imbibition rate was observed after the initial stage of exposure with the water bath. For this reason, the water content profiles reported here were collected in the first ~13,310 s (3.7 h) of contact for comparison.

Let us focus our attention on the effect that the artificial weathering exerts on the water absorption behavior in our samples. Concerning samples exposed to thermal and RH stresses, namely, PA1 and PA4, a significant difference in the amount of absorbed water can be clearly observed. In fact, whereas, in PA4, the WC value was found to be equal to ~3% (with a strong increase to ~5% between ~25 and ~35 mm), in PA1, WC is remarkably reduced, reaching approximatively ~0.8%. It is well known that the majority of the structural alteration induced by thermal/RH stresses in porous materials, in terms of both pore size distribution and degree of connection, must be ascribed to the thenardite–mirabilite phase transition [55]. As a consequence, an expansion of the initial microcracks of the stone should be expected, leading to visible alteration in the water absorption properties with respect to the reference sample. However, in the case of PA1, the presence of NS results in an overall diminishing of the water absorption tendency of the material, also limiting the effect on the pore network induced by artificial microclimatic stimuli. On the other hand, since NL does not occlude pores with the same efficiency as NS, the effect of thermal/RH stresses on the structure is more pronounced, as can be seen from the different “trend” of the corresponding water content profiles, with respect to those calculated in PA-TQ, as well as the mean water content values. More in detail, whereas, in PA-TQ, the presence of sharp “peaks” (especially between ~20 and ~50 mm) ascribed to a local increase in water content is...
observed, in PA4, a slight enlargement of such features can be recognized (see Figure 7). In the case of PA-TQ, the sharp jagged profiles reasonably reflect the natural pore size heterogeneity of the Pietra d’Aspra limestone, as extensively evidenced by the analysis of the pore distribution (see Figure 4).

On the other hand, the observed broadening of the localized WC peaks in PA4 suggests the presence of larger cavities/pores as a consequence of the thermally activated thenardite–mirabilite phase transition. As a result, the presence of slightly larger water-filled areas is expected, resulting in the observation of neutron radiographs characterized by extended water-rich domains. It is worth noting that the changes in pore dimensions as a consequence of the aforementioned phase transition are relatively small. However, as can be seen in Figure 8, where the comparison between the WC profiles in PA-TQ and PA4 is reported, the high spatial accuracy achieved by neutron radiography, in terms of liquid distribution within the stone network, allowed us to hypothesize such structural variations otherwise impossible to be highlighted through laboratory-based methodologies.

![Figure 8](image-url)  
**Figure 8.** Water content profiles for samples PA4 (NL consolidated) and PA-TQ (reference) between 20 and 50 mm. Dotted lines mark peaks which reveal, better than others, the enlargement induced by the thenardite–mirabilite phase transition activated by microclimatic stimuli (T/RH jumps).

Finally, the relatively high WC value (~4–5%) between ~25 and ~35 mm in PA4 was probably due to an extended fracture already present inside the stone or caused by the development of high crystallization pressures.

As far as PA2 and PA5 are concerned, accounting for samples exposed to 15 cycles of salt crystallization, a substantial reduction in the amount of absorbed water can also be detected. In PA2, WC was found to be equal to 1 wt.% after 3.7 h, reaching approximately ~18–20 mm. On the contrary, a remarkable increase in WC can be observed in PA5, with a mean value around 3–3.5%. This occurrence, in agreement with what was already observed in the case of PA1 and PA4 samples, supports the hypothesis that NS, more than NL, clearly enhances the structural properties of the analyzed stone by improving its overall hydrophobicity.

Salt crystallization cycles may lead to the formation of fractures and plane distortions to a major extent with respect to those induced by thermal/RH stresses. Accordingly, liquid solutions penetrate more efficiently and faster thanks to the presence of new “channels”, making the structure more susceptible to the presence of water even for short exposure times. In fact, by comparing the WC profiles of PA5 and PA-TQ, at equal contact times, we found that the mean water content value (wt.%) is always greater in PA5 than in PA-TQ. This occurrence suggests a scenario according to which the new artificially induced cracks in PA5, after 15 cycles of salt crystallization, are now filled with water through spontaneous imbibition, resulting in a higher WC value (~3.5%) with respect to that of PA-TQ (~2.5%). This hypothesis is also supported by the fact that, in PA6 (sample treated with nanolime...
and not subjected to any accelerated weathering tests), the water content is slightly lower (~2.5%) with respect to PA5, suggesting that the main reason for the observed increase in WC in PA5 must be ascribed to the salt crystallization test. NL, in this case, does not affect the limestone almost at all. In the case of PA1, the NS consolidant reduces the access of water through the occlusion of the outer shell pores, reducing the amount of included water and slowing the water dynamics with respect to both PA5 and reference (PA-TQ).

To further investigate the effects of the application of consolidants onto the limestone surface, the mean penetration depth of water ($h$) was evaluated as the sample height associated with the inflexion point (IP) of the decreasing WC profiles depicted in Figure 7.

IP can be considered as the position from the bottom of the sample at which a water-rich/dry zone transition occurs, and it is, hence, equivalent to the “front” of the wetted region during the capillary suction process. It is worth noting that an exact determination of IPs starting from neutron radiographs can be carried out easily only in those cases in which a uniform capillary absorption occurs, as observed in a large variety of porous materials characterized by a homogeneously distributed porosity [56,57]. In our case, however, the intrinsic heterogeneities, in terms of both natural constituents and pore size distribution, make the WC profiles not uniform; consequently, the evaluation of the IPs difficult to carry out. On the basis of these considerations, the IPs of the decreasing curves reported in Figure 7 and, hence, the position of the wetting front were approximated to the points corresponding to half of the maximum water content values, which are different in each investigated sample [58].

From a theoretical point of view, the behavior of $h$ as a function of time can be predicted by using the analytical Lucas-Washburn (L-W) equation (Equation (6)) [59,60].

$$h(t) = K \sqrt{t},$$

where $K$ is a coefficient, known as sorptivity ($m/s^{0.5}$), accounting for the tendency of a porous material to absorb/transmit liquid through natural imbibition. Generally, the L–W model assumes not only a complete saturation of the region behind the wetted front, but also a uniform pore size distribution of the material. Consequently, in materials characterized by a broad distribution of pores sizes, as in our case, deviation from the aforementioned model is often achieved. Nevertheless, by looking at the way in which the experimental values diverge from the expected ones, several considerations can still be made. In Figure 9, we report the $h(t)$ vs. $\sqrt{t}$ plot for all the investigated PA samples in the 0–13,310 s time range.

The comparison of the $h(t)$ vs. $\sqrt{t}$ plots for the PA-TQ specimen with those calculated for the treated limestones not subjected to any artificial weathering test (PA3 and PA6) indicates that the impregnation of the stone surface with NS visibly modifies the water transfer properties of the limestone to a major extent compared to the treatment with NL, both at the early and, mostly, at the late stage of the absorption test. This was also qualitatively confirmed by the extremely slow water kinetics observed in PA1, PA2, and PA3 with respect to that observed in PA4, PA5, and PA6.

The observed trends of $h(t)$ vs. $\sqrt{t}$ strongly deviate from the L-W model, especially for $\sqrt{t} > \sim 30 s^{0.5}$, indicating a heterogeneity of the porous medium in which the capillary rise of water is due to the contribution of several pore families with different size distribution. Nevertheless, below $\sqrt{t} \sim 30 s^{0.5}$, a diffusion-like behavior, as that predicted by the L-W model, can be reasonably hypothesized. This evidence can be justified thinking that, in the early stage of the test, the water motion is affected by the presence of a pore’s distribution influenced by both the stone and the consolidating agents, limited to the penetrated thickness. Accordingly, the occlusion of pores having different dimensions (ranging from $\sim 0.01$ to $\sim 100 \mu m$) leads the system from its natural widespread heterogeneity condition (see Figure 4) toward a more uniform one, with several pore families of different sizes partially or completely obstructed.

Although the quantitative evaluation of the sorptivity $K$ would not be reliable, since neither a full saturation of the region behind the water front (see Figure 6) nor a uniform pore network during all the imbibing process are satisfied, several considerations from Figure 9 can still be made.
The experimental evaluation of $h$ was corrected by taking into account the initial thickness of manually added water. Accordingly, all the inflexion point (IP) positions were shifted of 1-3 mm according to the initial thickness of water contained in the aluminum basement, whose contribution can be clearly seen as the highest peaks in the low-millimeter range in Figure 7 for PA2, PA5, PA6, and PA-TQ. In this way, a comparison of all the calculated $h$ values with respect to a common reference can be carried out with high accuracy. In black, the evolution of the water front position for PA-TQ is displayed, whereas the blue and red curves account for the NS- and NL-treated samples, respectively.

Concerning samples PA1, PA2, and PA3, the application of NS slows the water rise, which appears extremely hindered for both short and long exposure times (see blue curves of Figure 9). This occurrence testifies to a massive occlusion of pores preventing the water to flow upward (all three samples remain stacked below 20 mm), and then reducing the risk of alveolar decohesion, erosion, and the inclusion of damaging substances (with the mean water content also being extremely low; see Figure 7). This confirms, again, that the use of Nano Estel® (nano-SiO$_2$) as a consolidating agent against artificial weathering is much more convenient with respect to CaLoSil® (nanolime). Finally, it is worth noting that changes in the water kinetics between all PA samples can also be ascribed to the different penetration depth of the consolidating product within the porous structure. It is reasonable to assume that, according to the amount of penetrated NS or NL, the water diffusion behavior may also vary, leading to slight alterations of the natural imbibition phenomenon.

4. Conclusions

In this work, a neutronographic investigation was carried out on Pietra d’Aspra limestone, a particular type of lithotype extensively used in Baroque buildings of Palermo city (Sicily, Italy). The paper focused on the evaluation of the effectiveness of two different commercially available consolidants, i.e., nanosilica (NS) and nanolime (NL), against naturally occurring degradation
phenomena (i.e., temperature/RH jumps and salt crystallization), to which buildings and monuments are continuously exposed. The petrographic and physicochemical properties of the investigated limestone were firstly assessed using standard laboratory-based methodologies. Results from petrographic and petrophysical investigations on Pietra d’Aspra biocalcarenites highlighted the heterogeneity of pore network systems in these types of rocks. These features are particularly interesting parameters for interpreting damage processes and their role in the physical and mechanical behavior of the stone. Then, the collected neutron radiographs allowed us to gain insight into the water absorption behavior exhibited by the Pietra d’Aspra stone, revealing a strong relationship between the pore characteristics, in terms of both size and distribution, and the occurring natural water imbibition. In particular, the analysis of the maximum amount of water content and height of the water front revealed a better performance of NS with respect to NL. Furthermore, the effect of several accelerated artificial stimuli on the fluid rises by capillarity was also successfully highlighted through neutron radiography, which, thanks to its high spatial resolution, allowed us to experimentally observe localized water quantities, as well as inner stone alterations (i.e., enlargement of pores following artificially activated phase transitions), otherwise impossible to be revealed. The performed studies of petrophysical properties and capillary imbibition kinetics are indispensable for understanding the degradation mechanisms of Pietra d’Aspra, in order to give an appropriate response for future restoration interventions.

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