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Influence of the intrinsic characteristics of mortars on their biofouling by pigmented organisms: Comparison between laboratory and field-scale experiments

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Abstract

Biodeterioration of mortars by the photosynthetic microorganisms is affected by their intrinsic properties such as porosity, roughness and surface pH. The influence of these parameters was examined using an accelerated fouling test in laboratory and a natural fouling test in the real-world (\textit{in-situ}). Basing on color measurement and image analysis, the impact of each intrinsic parameter was evaluated. The results differed from a scale to the other one.

No influence of porosity was measured on the algal colonization rate in the laboratory test whereas, a high porosity seemed to increase slightly the bioreceptivity of the mortars exposed outdoor. The roughness, in both tests, promoted the microbial colonization. However, the discrimination of roughness grades was better in the laboratory test than in the \textit{in-situ} one.
The surface pH influenced remarkably on the accelerated biofouling test but not on the *in-situ* one. These dissimilarities resulted from the differences in experimental configurations of the two tests.

**Keywords**: mortar, biofouling, intrinsic properties, laboratory test, *in-situ* test

1. **Introduction**

The colonization of building facades by microorganisms is a subject that concerns more and more the civil engineering community. Indeed, this phenomenon changes the aesthetical appearance of materials and, in later stage, can even compromise the durability of structures by corrosion or by physical degradation induced by the microorganism (Maury-Ramirez et al., 2013). The type of microorganisms colonizing cement based building facades is diverse. But the analyses of *in-situ* samples show that algae are one of the initial and main colonizers (Gaylarde and Gaylarde, 2005). These microorganisms allow the successive implantation and the growth of other biological organisms such as lichens and bryophytes. If no solution is found, the pteridophytes and higher plants may also appear (Perrichet, 1984; Deruelle, 1991; Barberousse, 2006). The microbial development must be eliminated to avoid worse consequences to buildings. The more the cleaning maintenance activities are repeated the more expensive the cost.

Several factors such as the global climate, the local environment, the building design and the facade materials influence the biological fouling.

The rainfall, the wind and the temperature consist in the climatic parameters influencing together the water availability of facade materials which is an essential element to the microbial metabolism. A permanently wet facade promotes the growth of algae and higher
plants (Ariño and Saiz-Jimenez, 1996; Loh, 2002). Therefore, the facades represent a higher susceptibility to biofouling in the rainy regions and during the heavy rainy season (Young 1997). However, a high temperature induces water evaporation by heating the materials. Similarly, the wind can cause a drying phenomenon.

The local environmental conditions, such as topography and nature of the ground, presence of moisture and/or industrial activities around the buildings, should be taken into account. As an example, a house roof, located close to seaside, can be totally colonized by nitrophilous lichens, characterized by the yellow or orange, in a few years (Deruelle, 1991). Indeed, sea sprays, by providing nitrogen to the building surface, favor the microorganism growth. In addition, the humid environments set near a lake, a river, trees or shrubs, are favorable to the appearance of stains on the walls.

The growth of biological fouling is also depending on the orientation of the facade. Indeed, the north-facing facades which are wetter and less sunny, get colonize faster (Ariño and Saiz-Jimenez, 1996; Young, 1997; Barberousse, 2006). Similarly, a facade exposed to dominant winds seems to be colonized more easily than the other sides of the same building. The wind can transport both the rain and the biological propagules at the facades promoting the biofouling.

Parts of building often moistened for long periods, or easily covered of propagules, are highly sensitive to the biological colonization (Wee and Lee, 1980). The biofouling often grows at the foot of walls, the junction of different coatings and the overhanging elements (cornices, moldings, balconies, etc.) (Wee and Lee, 1980; CSTB, 2005; Barberousse, 2006).

In addition to external conditions, the biological development is affected by the intrinsic characteristics of the facade, defined as its bioreceptivity (Guillitte, 1995). These
characteristics can be divided into physical properties (porosity, roughness and hydrodynamic properties) and chemical ones (chemical composition, pH surface).

Since the last thirty years, several researches have been devoted to the study of the building-materials bioreceptivity. In all these studies, the experimental tests were conducted with more or less accelerated methods of the biological growth on materials. The goal of these accelerated methods was to reduce the test duration, which normally required many years in the real-world. The experimental parameters were thus optimized to favor the growth of microorganisms (Dubosc et al., 2001; Barberousse, 2006). Different methodologies, concerning the preparation of materials, the process of inoculation and humidification, the temperature and the photoperiod of the test were developed (Grant and Bravery, 1985; Guillitte and Dreesen, 1995; Ohshima et al., 1999; Dubosc et al., 2001; Miller et al., 2006; Miller et al., 2009; De Muynck et al., 2009; Tran et al., 2012). According to all these studies, the roughness of materials appears as one of the most important parameters. Indeed, the colonization of the building materials by microorganisms is promoted by the roughness of the surface. In fact the roughness provides many asperities and thus increases the physical anchorage of these micro-organisms (Dubosc, 2000; Tran et al., 2012). This effect of roughness on the biofouling was confirmed by observations of buildings in real conditions (Wee and Lee, 1980; Darlington, 1981; Pietrini et al., 1985; Joshi and Mukundan, 1997). The total porosity, the pore size distribution and the porous network, should be taken into account, because the absorption and the water retention of materials are controlled by these characteristics (Warscheid et al., 1993; Ohshima et al., 1999; Crispim et al., 2003; Miller et al., 2006; Miller et al., 2009). Furthermore, the inhibition of the algal growth by high pH of material surface is commonly mentioned (Grant, 1982).

Most of previous studies propose only a qualitative evaluation of the biofouling or a comparison of very different materials. In order to accelerate the fouling, mortars are usually
aged (by carbonation and/or leaching) to decrease the surface alkalinity. But the effect of the accelerated aging is rarely evaluated. Moreover, the experimental conditions of accelerated tests are significantly far from the real conditions. The investigations in field-scale are rarely conducted because of the long duration of the experiments (Young, 1997; Ohshima et al., 1999; Lengsfeld and Krus, 2004). Thus, the relationship between bench-scale experimental results and field-scale experimental results is scant.

This work aims to clarify the effect of porosity, roughness and carbonation, on the building-materials colonization by algae at laboratory and in-situ scale. One unique mortar formulation has been investigated. The kinetic of the biological colonization was studied.

2. Materials and methods

2.1. Preparation and characterization of samples

Portland cement (CEM I 52.5, Holcim), siliceous sand (Sibelco DU 0.1/0.35), calcareous filler (Omya) and cellulose ether (Hydroxylethyl Methyl Cellulose, SE-Tylose) were mixed according to the proportions summarized in Table 1 to prepare the studied samples. The water-to-cement ratio ($w/c$ (wt/wt)) was fixed at 1 and 1.2. These two $w/c$ were considered in order to modify the porosity without changing the chemical composition of mortars.

Table 1

<table>
<thead>
<tr>
<th>Component</th>
<th>Cement</th>
<th>Sand</th>
<th>Calcareous Filler</th>
<th>Admixture$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>% mass of dry mixture</td>
<td>30</td>
<td>65</td>
<td>5</td>
<td>0.27</td>
</tr>
</tbody>
</table>

$^a$ in addition to dry mixture (cement, sand and filler)
The variation of sample surface pH was achieved by a curing process. Mortars were subjected (called carbonated mortars) or not (called uncarbonated mortars) to ageing by accelerated carbonation prior to the start of biofouling test.

After mixing, the fresh mortar was casted into 50 cm × 50 cm × 1 cm extruded polystyrene molds. The fresh mortar was stored at 21 ± 1 °C and 95 ± 5 % of relative humidity during 28 days for the uncarbonated samples. The mortar plates were then cut into samples of 20 cm × 8 cm × 1 cm and of 20 cm × 30 cm × 1 cm for the laboratory tests and the in-situ tests, respectively. For the carbonated samples, the fresh mortar was stored in the aforementioned conditions for only 7 days before being cut. This premature curing end aimed to shorten the total duration of specimen production and to obtain the closest total storage time as possible between carbonated and uncarbonated samples.

The samples were then exposed to pure CO₂ at 21 ± 1 °C and 65 ± 5 % of relative humidity until the complete carbonation. This carbonation process required 36 days. The pH was verified by spraying a phenolphthalein solution (0.2 % in ethanol) on the entire thickness of specimen. The discoloration of the solution indicated a diminution of pH to about 9 (Thiery, 2005).

Moreover, the surface pH of mortar was measured by means of a surface electrode (WTW Sentix Sur). For each sample, six measurements were performed at six positions distributed over the sample surface. A constant volume of distilled water (0.1 ml) was deposed on the related position. The electrode was kept in contact with the sample surface during a fixed time of 30 seconds. The pH value was then recorded and the surface pH of sample was the average of these six values. The surface pH of sample was monitored twice a week during the carbonation process.
The pore size distribution and the total porosity of materials were determined using mercury intrusion porosimetry (Micromeritics Autopore IV 9400). Three samples were measured for each mortar formulation, i.e. w/c ratio of 1 and 1.2, both uncarbonated after 28 days of curing and carbonated after 7 days of curing and 36 days of accelerated carbonation. They were beforehand dried by acetone to evacuate the open pores.

To obtain several surface roughnesses, three surface finishing methods were applied during the sample setting. The first one consisted in smoothing the surface of fresh mortars with a ruler and the two others in scratching the surface of the setting mortars with sponges of two different roughnesses.

The roughness of the mortar specimens was evaluated by means of an optical profilometer (CHR-150-L). The arithmetic average of the height \( R_a \) was determined from the measurements of the surface profile (Gadelmawla et al., 2002). Measurements consisted in analyze 161 profiles of 7 cm length and spaced of 500 \( \mu \)m. The data was recorded each 20 \( \mu \)m along each profile. The value \( R_a \), representative of the sample roughness, corresponded to the average of 563.661 measuring points.

Each mortar was labeled by four codes. The first one corresponds to the w/c ratio, the second to the carbonation state, the third to the roughness and the last to the scale of the biofouling test. For example, a sample labeled 1UC-R1-Lab corresponded to the uncarbonated mortar prepared with a w/c ratio of 1 with the smoothest roughness and tested in the laboratory experiment.

2.2. Lab-scale and field-scale biofouling tests

The laboratory experiments consisted in an accelerated algal growth on cementitious materials. A closed device containing an algal suspension was set up. The microalgae *Klebsormidium flaccidum* was selected in this research basing on its representativeness and
ease of liquid culture. The suspension was periodically sprinkled on the samples surface. The
run-off period was fixed to 90 min every 12 h. The light was provided by two neon lamps
during 12 h per day, and was set to start simultaneously with the run-off cycles. All the
experimental configurations (rate of sprinkling, light intensity, temperature, initial
concentration of the algal suspension and angle inclination of the samples) were kept
constant. The details of the experimental approach were described in previous papers (Tran et
al., 2012; Tran et al., 2013).

In laboratory experiments, specimens with dimension of 20 cm × 8 cm × 1 cm were tested
in triplicate for each material. Carbonated and uncarbonated samples were tested separately.
Therefore, 18 samples were introduced in the test-chamber for each experiment.

The field-scale experiments consisted in a natural development of microorganisms on
cementitious materials. The mortar samples were placed on a stainless steel frame in a private
green park close to Grenoble, France. The sample dimensions were larger than the ones used
in the laboratory test, i.e. 20 cm × 30 cm × 1 cm. The larger dimensions aimed to minimize all
the non-representative effect resulted from the natural incidental events such as a random
intense inoculation of microorganisms at an individual sample points, the deposition of bird
droppings on the sample surface, etc.

The exposure of materials to the outdoor conditions started on 26 June 2009 and lasted one
year and a half. The experimental configurations were chosen in order to favor the biological
colonization of the sample surface. The samples were placed near trees, facing the north
direction and inclined with an angle of 45 °. The specimens were arranged in two rows of
which the first was located at 1 m above the ground to avoid splashing during rainy periods.
The contamination between specimens by water flow was also avoided (Fig. 1).
Each material was tested in triplicates. The mortars with a w/c ratio of 1 were studied at carbonated and uncarbonated states, with the three roughnesses. For the samples with a w/c ratio equal to 1.2, only the carbonated mortars with the two highest roughnesses were examined.

2.3. Evaluation criteria of biofouling

The biological colonization of mortar surface was evaluated by means of image analysis and colorimetric measurements.

Concerning the image analysis, the surface of samples was digitized using Epson V300 office scanner. The images obtained in the RGB color space were converted into the YIQ color space. The distinction between the areas covered by algae and the surfaces of clean mortars were achieved on the Q channel (Tran et al., 2012). The colonization rate ($X(t) (%)$) was then given by the ratio of the fouled areas to the total surface of the sample. The samples from the lab-scale and field-scale experiments were scanned every weekday and at least every two months, respectively.

The colorimetric measurements were performed using colorimeters (Minolta CM-2600d for the laboratory tests and Konica Minolta Chroma CR-410 for the in-situ tests). For each
specimen, the measurements were conducted at fixed positions spread over the whole sample surface. Thirty six and eighteen measurements were monitored on a lab sample and an *in-situ* one, respectively. The sample area of each measurement was of 0.5 cm$^2$ and 19.6 cm$^2$, respectively. The CIE Lab color space data ($L^*$, $a^*$, $b^*$ coordinates) were collected. The data was monitored three times a week on the lab specimens and at least every two months for the field-scale specimens. In order to evaluate the color changes of the test specimens, for each color coordinate ($L^*$, $a^*$, $b^*$), the difference between sample at a given time and the same sample at the initial time (uncolonized sample) was determined ($\Delta L^*$, $\Delta a^*$, $\Delta b^*$).

### 3. Results

#### 3.1. Characterization of materials

The porosity, the surface pH and the roughness of each mortar are given in Table 2. For the samples made with $w/c$ ratio equal to 1 (i.e. uncarbonated and carbonated), the values of the porosity, the surface pH and the roughness R1 were those of a previous paper (Tran et al., 2012).

#### Table 2

Characteristics of mortars

<table>
<thead>
<tr>
<th>Ratio w/c</th>
<th>Porosity (%)</th>
<th>Surface pH</th>
<th>Code</th>
<th>$R_s$ ($\mu$m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Uncarbonated 1</td>
<td>37.2 ± 0.1</td>
<td>11 ± 0.4</td>
<td>Roughness 1</td>
<td>1UC-R1</td>
</tr>
<tr>
<td></td>
<td>1.2</td>
<td>38.9 ± 0.3</td>
<td>Roughness 1</td>
<td>1.2UC-R1</td>
</tr>
<tr>
<td>Carbonated 1</td>
<td>32.1 ± 1.9</td>
<td>9.0 ± 0.1</td>
<td>Roughness 1</td>
<td>1C-R1</td>
</tr>
<tr>
<td></td>
<td>1.2</td>
<td>36.2 ± 0.1</td>
<td>Roughness 1</td>
<td>1.2C-R1</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Roughness 2</td>
<td>1.2C-R2</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Roughness 3</td>
<td>1.2C-R3</td>
</tr>
</tbody>
</table>
The total porosity of uncarbonated mortars varied from 37 to 39 %, which is coherent with the literature (Monge 2007). The mortars with the highest w/c ratio were slightly more porous. According to Lafhaj et al. (2006), the porosity increases non-linearly with the w/c ratio. When this w/c ratio is high, its effect on the total porosity is attenuated.

The carbonated mortars were less porous than the uncarbonated ones, i.e. 32 % for 1C against 37 % for 1UC and 36 % for 1.2C against 39 % for 1.2UC. It is known that the carbonation leads to a reduction of pore volume. Houst (1992) noted that this reduction of the total porosity is even more enhanced by a low w/c ratio.

Depending on the curing process, two levels of surface pH were obtained. The surface pH reached 11 with the standard curing (uncarbonated samples) and 9 by accelerated carbonation. Moreover, no effect of the w/c ratio on the surface pH was detected. Indeed, the same value of surface pH was measured for both w/c ratios.

Three levels of roughness were obtained whatever the curing process and the w/c ratio. The smoothest surface (R1), without reliefs, was achieved by the method of finishing to the rule. The rougher surfaces were obtained by scratching the sample surfaces with sponges. Both strong reliefs and asperities appeared on the roughest surface (R3). Moreover, a remarkable discrimination between the highest roughness (R3) and the intermediate one (R2) was observed. In fact, the $R_a$ values were 2.5 to 5 times higher for R3 than for R2. As a comparison, the difference between the R2 and R1 was only about 1.5 times.

3.2. Biological colonization of cementitious materials in laboratory and field test

Fig. 2 illustrates the biological fouling of specimens investigated in the accelerated tests and the in-situ experiments over time.
**Fig. 2.** Surface colonized by algae over time for carbonated mortars, w/c of 1.2 and roughness R3.
In the laboratory tests, the colonization of the surface by *Klebsormidium flaccidum* created a dense and velvety mat as previously shown by Tran et al. (2012; 2013). The first algal spots appeared at privileged sites of the surface, such as air bubble holes or asperities formed by the roughness. The extension of fouling resulted from the growth of the existent spots and the adhesion of new ones. On the sample surface of lowest roughnesses (R1 and R2), the algal development formed streaks due to the suspension flow. This fouling form was usually observed on the building facades. On the roughest mortars (R3), the fouling followed the asperities on the surface which conducted the flows of the algal suspension.

The sample exposure in the natural conditions began at summer (end of June 2009). The first settlement appeared on the mortars 1.2C-R3, after 11.6 months of exposure. The colonization rate by microorganisms of this mortar surface is shown in the Fig. 3. Following the latency period, the biological colonization developed rapidly. Four and a half months later, the fouling was observed on the whole *in-situ* specimens.

![Fig. 3](image)

**Fig. 3.** Colonization rate for carbonated mortars, w/c of 1.2 and roughness R3 in the *in-situ* test.

The same latency time was observed by Dubosc (2000) and Young (1997) for carbonated mortars and sandstone samples, respectively. The biological development accelerated during the autumn period (August to November, i.e. 13 to 17 months of exposure). The same
observation was made by Warscheid et al. (1993) on stone samples exposed to outdoor conditions in different locations in Germany.

The propagules of microorganisms might be already settled on the mortar surface since the spring. Fig. 4 shows the monthly average values of rainfall and air temperature during the experimental time. In spring 2010 (April and May corresponding to 10 and 11 months respectively), the meteorological data highlighted high precipitation and medium temperature (i.e. about 15 °C). The following summer corresponded to the harsher climate, especially in July (12-13 months), with the highest average temperatures and the lowest precipitations of the year. During this period, the water available for the microorganism development was very limited. Then mild temperatures accompanied with a suitable rainfall in autumn (September and October corresponding to 15 and 16 months of exposure respectively) favored remarkably the algal growth. From December (18 months), the detachment of algae from the samples surface was observed. It corresponds to the winter period characterized by very low temperatures, weak density of lighting and snow. The aging of microorganisms in this season might be the reason of theirs detachment.

Fig. 4. Monthly average air temperature and cumulative precipitation during the *in-situ* test.

Fig. 5 represents the color changes of carbonated mortars of roughness R3, tested in laboratory and field-scale. All the graphs show the mean values and the standard deviation of
three specimens. The evolution of clarity $\Delta L^*$ and chromaticities $\Delta a^*$, $\Delta b^*$ showed the same
trends for the two test scales. The $\Delta L^*$, $\Delta a^*$ decreased while the $\Delta b^*$ increased with time. The
biological colonization led to more and more dark ($\Delta L^*$), green ($\Delta a^*$) and yellow ($\Delta b^*$)
sample surface. This color is characteristic of green algae. The main colonizers on the in-situ
samples could be attributed to these microorganisms.

![Graphs showing the evolution of $\Delta L^*$, $\Delta a^*$, and $\Delta b^*$ over time.]

**Fig. 5.** Evolution of the clarity $\Delta L^*$ and the chromaticities $\Delta a^*$, $\Delta b^*$ of mortars with the
roughness R3 in lab- and field-scale tests.

### 3.2.1. Effect of w/c ratio

Fig. 6 shows the influence of the w/c ratio (1 and 1.2) on the biofouling rate of carbonated
mortars (three roughnesses).
In the lab test, the algal colonization curve was of sigmoid type. Three steps can be identified: a latency step, an exponential growth step and a step of stagnation. For each roughness (R1, R2 and R3), no effect of the w/c ratio was identified on algal coverage. In fact, for the roughest mortars (R3), the fouling rate evolved quite identically. The algal colonization appeared after 6 testing days and the entire surface was covered after 30 days. For the two other roughnesses, with such experimental standard deviation, the colonization rate of the less porous mortars (1C) could be considered similar to the one of the more porous mortars (1.2C).

Unlike in the accelerated test, in the real conditions, an effect of the w/c ratio was detected. Indeed, the mortars made with a w/c ratio of 1.2 presented a faster microbial development
than the ones prepared with a w/c ratio of 1. For the 1.2C-R3 samples, the biofouling appeared between 11 and 13 months of exposure. At the same moment, no algal colonization was observed on the mortar 1C. The following measures, realized two months later, showed a coverage area of 80 % for 1.2C mortars against only 30 % for 1C mortars. For the intermediate roughness (R2), the effect of w/c ratio was less important. 30 % of the mortars 1C was colonized while it was 2 times higher on the mortar 1.2C. One month later, only the more porous mortar was totally colonized.

The color changes of the roughest mortar confirmed the results obtained from the images analyses (Fig. 5). Indeed, the color intensity of 1.2C-R3-Field mortars evolved faster than the one of 1C-R3-Field mortars. Furthermore, the change in color intensity between these two mortars occurred after 13 months of exposure. Concerning the bench-scale experiments, the evolution of the color and the colonization rate coincided perfectly regardless of the w/c ratio. Indeed, an increase in the color intensity was detected between 4 and 28 days of exposure. Then, no color evolution was observed since the 28th days of testing, corresponding to the necessary time for a complete biofouling.

3.2.2. Effect of the roughness

For the bench-scale test, the biofouling rate was obviously influenced by the surface roughness of samples (Fig. 6a). The effect of the three levels of roughness tested on the biofouling were quite distinct, whatever the w/c ratio (1 and 1.2). The smoothest roughness (R1) exhibited the lowest colonization rate and the highest latency time. Indeed, the latency time was around 18 days of testing for roughness R1, against 9 and 6 days for roughness R2 and R3, respectively. Similarly, the complete colonization was achieved more quickly on the rougher mortars, i.e. in 30 days for R3, against 51 and 56 days for R2 and R1, respectively. Furthermore, the curve slope evolved in the same way as the roughness during the
exponential growth phase. The more the roughness was high, the more the colonization was quick.

The roughness impacted also the biological development of mortars exposed in the real-world (Fig. 6b). However, unlike bench-scale test, only two grades of roughness (i.e. rough and smooth) were observed. The mortars of very different roughness, R2 and R3, showed no difference in bioreceptivity to visible photosynthetic organisms. This result was particularly obvious for w/c = 1. Conversely, a transition from a smooth surface (R1) to a rough one (R2) remarkably increased the bioreceptivity of mortars. Indeed, the colonization appeared slower for the mortar R1 than for the two others. The entire surface of samples R2 and R3 was colonized after 17 months (November 2010). For the same duration, only 60 % of mortar surface R1 was colonized. Beyond this time, a decrease in the covered area was noticed for all samples. It has been assigned to detachment of algae. The detachment rate was similar for all mortars of w/c equal to 1. However, for mortar of w/c equal to 1.2, smooth surface showed a faster decrease in the coverage area compared to the rough one. Hence, the effect of roughness on the microbial detachment rate remains difficult to analyze.

3.2.3. Effect of the accelerated carbonation

Fig. 7 illustrated the evolution of the colonization rate for carbonated and uncarbonated mortars mixed with a w/c ratio of 1, for the lowest and the highest roughness.
The carbonation seemed to be the most decisive parameter on the bioreceptivity of materials for the accelerated fouling test. It significantly shortened the latency time, and accelerated the rate of algal colonization. As showed in the Fig. 7, the latency time of 1C-R1 mortar was of 18 days. This value was 2.6 times less than that of 1UC-R1 mortar (46 days). Similarly, the colonization of 1C-R3 mortar began after 7 days against 26 days for 1UC-R3 mortar. The complete colonization was reached in 30 days for the 1C-R3 mortar against 51 days for the 1UC-R3 one. It is worth noting that, the total colonization of 1UC-R1 was not obtained because of a precocious stop of testing. Nevertheless, basing on the roughest mortars, the slope of the curve was greater for carbonated mortars than for uncarbonated ones.

In the field-scale test, it seems that the accelerated carbonation did not result in different growth profiles. The biofouling evolved quite identically on carbonated and uncarbonated mortars. For both mortars 1UC and 1C, and of roughness R1 or R3, the colonization by microorganisms began at the same moment, i.e. after 13 months. Then, 4 months later (November, corresponding to 17 months of exposure), the entire surface of all rough samples (R3) was colonized. At this same time, the algal colonization reached a common rate of around 60% for smoothest samples (R1), carbonated or uncarbonated. From 18 months...
(December 2010), a decrease of fouled area was observed on the entire surface due to the microbial detachment. It seems that the detachment rate was not dependent on the initial carbonation state of mortars.

4. Discussion

The results obtained from the color measurements and the image analysis, are coherent and complementary. Indeed, the effect of intrinsic characteristics of mortars on their bioreceptivity was found identical by these two methods. Due to monitoring the entire surface of samples, the color changes are well representative of the fouled area.

The color evolution of the surface of the samples, for both laboratory and in-situ tests, was quite identical, i.e. more and more dark, green and yellow. The fouling of the in situ samples is thus presumably associated to green algae. These microbial species are the primary and main colonizers on building facades as noticed by several authors (Wee and Lee, 1980; Grant, 1982; Ortega - Calvo et al., 1995; Tomaselli et al., 2000; Barberousse, 2006).

The influence of w/c ratio on the accelerated fouling was negligible. The reason is related to experimental conditions rather than to modest difference in porosity between two studied mortars. Indeed, the relative humidity in the laboratory test chamber is permanently very high (i.e. from 80 to 100 %) (Fig. 8). In this condition, the water content into the samples is always abundant and so is sufficient to the algal development.
In order to verify this assumption, the filling rate of pore by water ($\varepsilon$ (%)) was calculated
(Dubosc, 2000). It corresponds to $q/p$ ratio, where $p$ (%) is the sample porosity, and $q$ (%) is
the amount of water absorbed by a unit of sample volume, calculated as follows:

$$q(\%) = 100 \times \frac{V_{\text{water}}}{V_{\text{sample}}} = 100 \times \frac{m_{\text{water}}}{\rho_{\text{water}} \times V_{\text{sample}}}$$

$V_{\text{water}}$: volume of water absorbed by the sample (cm$^3$);

$V_{\text{sample}}$: apparent volume of the sample (cm$^3$), calculated from three dimensions of sample;

$m_{\text{water}}$: amount of water absorbed by the sample (g) that is equal to the difference between
the mass of sample in the humid condition and in the dry one.

$\rho_{\text{water}}$: density of water (1 g.cm$^{-3}$)

The humid sample weight was measured 6 times along the diurnal cycle and 2 times along
the nocturnal one. Before the start of accelerated fouling test, the sample was dried at 40 °C
until obtaining constant weight. This weight was considered as the mass of dry sample.

The amount of water absorbed by the sample reached a maximum value just after the
sprinkling cycle and remained constant during all the nocturnal period. During the diurnal
period, the water evaporated under the warming effect induced by neon lights. However, the

Fig. 8. Daily cycle of temperature and relative humidity in the test chamber.
amount of evaporated water was weak. Indeed, at the end of the diurnal period, the remaining water into the mortars filled at least 75 % and 60 % of the porosity of uncarbonated and carbonated samples, respectively.

The water, being an indispensable element for the biological metabolism, is thus no longer a limiting factor for the biological development. This observation was confirmed by previous studies regarding two mortars made with w/c ratios equal to 0.5 and 1 (Tran et al., 2012; 2013). Despite of the considerable difference in porosity, these mortars exhibited completely identical bioreceptivity.

In natural condition, the impact of the w/c ratio on the colonization rate of samples was exhibited. The microbial fouling seemed to be favored by the highest w/c ratio (1.2).

To evaluate the porous network of mortars, analyses were performed after 17 months of exposure to the outdoor conditions. PIM’s results indicate a modification of the porous network. Indeed, the total porosity of the 1C-R3 and 1.2C-R3 mortars increased from 32 to 36 % and from 36 to 38 %, respectively. Moreover, the single peak characterizing the pore entrance diameter shifted to larger diameters (Fig. 9). Indeed, the pore inlet diameter, that was equal to 0.3 μm at the beginning of outdoor exposure, was about 0.5 μm at the end of the experiment for the 1C mortar. For the 1.2C mortar, the pore inlet diameter evolved from 0.75 to 1.6 μm. The enlargement of the pore entrance diameter could be due to the leaching effect by rain and to the activities of biological agent such as bacterial communities and/or fungi (Warscheid and Braams 2000). These results, which corresponded to the average values of the entire thickness of samples, might be amplified at the surface.
Fig. 9. Pore size distribution of carbonated mortars with a \( w/c \) ratio of 1 and 1.2 at the beginning and end of the \textit{in-situ} test.

The mortars made with a \( w/c \) ratio of 1.2 were more porous than those formulated with a \( w/c \) ratio of 1 and exhibited a greater bioreceptivity. The effect of porosity on the biofouling rate of samples exposed outdoor are in agreement with the literature (Warscheid et al., 1993; Ohshima et al., 1999; Crispim et al., 2003; Prieto and Silva, 2005; Miller et al., 2006; Miller et al., 2009).

Previous studies highlighted an important role of roughness on the biological colonization (Prieto and Silva, 2005; Miller et al., 2006; Miller et al., 2009; Tran et al., 2012; Tran et al., 2013). By providing asperities, the roughness favors the physical attachment of microorganisms which is dispersed by the water flow or the wind. Consequently, the rougher the surface is, earlier the biological fouling begins and faster the colonization is. The recent accelerated fouling experiments allow confirming the effect of roughness. Indeed, the three grades of roughness studied induced a discriminating effect on the colonization rate. The same results were noticed by Tran et al. (2012) on others mortars mixed with a \( w/c \) ratio of 0.5.

For \textit{in-situ} samples, the visible biofouling due to phototrophic organisms appeared approximately at the same moment whatever the roughness of mortars. Indeed, for all the roughnesses, the first spots were detected after about 13 and 15 months of exposure. Based on
the results of laboratory tests, the in-situ samples of the roughness R3 should have been firstly colonized, successively followed by mortars of the roughness R2 and then R1. However, in the in-situ test, the season, the external relative humidity and temperature could impact the colonization and thus hided the roughness effect.

In the natural conditions, only two grades of roughness were observed. The two roughest mortars showed the same bioreceptivity to pigmented organisms despite considerably different roughness. This tendency remains coherent with that obtained in laboratory test. In fact, a nonlinear relationship between roughness and algal fouling rate is noticed. A close bioreceptivity was observed between mortars of high roughness (Tran et al., 2012). The non-differentiation in the outdoor test could be induced by a natural inoculation much less intense than in accelerated tests.

Concerning the accelerated carbonation, this intrinsic characteristic of the samples appeared as the most important parameter which affected the biological growth in the accelerated fouling test. Unlike the effect of roughness which favors the ability of algae to physically cling to the surface, the pH and/or carbonation disturbs the biological metabolism. Carbonation leads to a decrease in the surface pH of samples and to a lower increase in pH of the algal suspension flowing on test specimens. Consequently, the K. flaccidum algal cells are in less alkaline conditions and thus in less stressful environment for their growth with carbonated samples than with uncarbonated ones. Indeed, K. flaccidum is known as an acidophilic microorganism. The ability of attaching and growth of algae on a surface are thus promoted by the carbonated of samples (Tran et al., 2012; 2013).

The growth of the algal spots on the sample surface is the consequence of the vegetative and cellular multiplication which is favored at low pH (Škaloud, 2006). The attachment of algae on the surface of the substrate is governed by adaptive metabolic interactions between
algal cells and the substrate (Fattom and Shilo, 1984; Finlay et al., 2002; Barberousse et al., 2006). The algal extracellular polysaccharides could play the role of glue (Robins et al., 1986; Gantar et al., 1995; Barberousse et al., 2006). These metabolites depend on the algal species considered and the substrate. They are involved in the initial contact between the cell and the surface, and act over time (Barberousse et al., 2006).

As a consequence, carbonated mortars are highly receptive to colonization by microorganisms. Algal colonization begins much earlier, and occurs faster compared to the uncarbonated mortars.

In the field-scale test, the influence of carbonation was not highlighted. Indeed, the biofouling of mortars, carbonated or uncarbonated, was detected at the same moment. Perhaps, the favorable climatic conditions and sufficient inoculation conditions were not met at the beginning of testing. In fact, the samples were exposed through the summer. Therefore, spring, the most favorable season for the microorganism spreading and their growth was past. Hence, the microbial colonization of carbonated samples was not initiated despite a favorable surface pH. After one year of exposure, the mortars were aged and weathered by leaching and natural carbonation. Consequently, the same surface pH (pH = 8) was measured for all the samples whatever their initial carbonation state. The bioreceptivity of mortars is thus identical when favorable conditions to the biological development have been in place. Therefore, the influence of the initial surface pH on biofouling is completely inhibited.

5. Conclusion

The present research has investigated the effect of the intrinsic parameters (porosity, roughness and carbonation) of mortars on their biofouling by photosynthetic organisms. This study was conducted by both laboratory accelerated tests and in-situ ones.
The impact of porosity on the biofouling of mortars is different for the two experimental test scales. This parameter has no effect on the biofouling rate in laboratory tests due to the experimental conditions. However, although more experiments of verification still required for the field-scale tests, high porosity seems to favor the biological colonization.

For the both test scales, the influence of the roughness is evidenced. A rough surface enhances the biological attachment. The discrimination of roughness grades was better in the accelerated tests than in the field-scale ones. This result could be explained by the fact that the accelerated test is performed in a closed circuit with an intense inoculation while the \textit{in-situ} test involves natural inoculation.

Thanks to its experimental configuration, the laboratory tests allow revealing the role of the accelerated carbonation. It is considered as a decisive parameter in the accelerated fouling. Indeed, carbonated mortars lead to a lower pH value of the mortar surface and the algal suspension in the test chamber. The algal development is thus significantly promoted and so the colonization rate is remarkably accelerated in the case of carbonated mortars. While in the natural conditions, due to numerous uncontrolled and random environmental factors, the effect of the carbonation was not observed. The evolution of biofouling is identical on the both carbonated and uncarbonated mortars. Due to the long exposure time and progressive ageing by lixiviation and carbonation, the surface pH, and thus the alkalinity, of carbonated and uncarbonated samples is the same.

The divergence of results between the laboratory and field-scale tests, prevent to correlate the two experimental scales. These dissimilarities could be remediated by improving the experimental protocol. In the accelerated test, a drying phase of samples between two sprinkling cycles could allow highlighting the role of porosity. A reasonable choice of the beginning of sample exposure to natural environment could highlight the effect of
carbonation. The effect of roughness in accelerated test could be closer than that in natural test by decreasing the inoculation intensity. This decrease in inoculation process could be achieved by reducing, either the initial concentration of the algal suspension or either the run-off.
References


