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Article Fabric-Reinforced Cementitious Matrix (FRCM) carbon yarns with different surface treatments embedded in a cementitious mortar: mechanical and durability studies

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Abstract: FRCM systems are nowadays increasingly used for strengthening and retrofitting of ex-9isting masonry and reinforced concrete structures. Their effectiveness strongly depends on the bond10that develops at the interface between multifilament yarns, which constitutes the reinforcing fabric,11and the inorganic matrix. It is well known that fabric yarns, especially when constituted by dry12carbon fibers, have a poor chemical-physical compatibility with inorganic matrices. For this reason,13many efforts are being concentrated on trying to improve the interface compatibility by using different surface treatments on multifilament yarns.15

In this paper, three different surface treatments have been considered, the first two consisting of yarns pre-impregnation with flexible epoxy resin or nano-silica coating while the third one is a process of fibers oxidation. Uniaxial tensile tests have been carried out on single carbon yarns to evaluate tensile strength, elastic modulus and ultimate strain, before and after surface treatments, and 19 also after yarns exposure to accelerated artificial aging conditions (1000 h in saline or alkaline solutions at 40 °C), to evaluate the long-term behavior in aggressive environments. 21

Pull-out tests on single carbon yarns embedded in a cementitious mortar were also carried out, under normal environmental conditions and after artificial exposure. Epoxy proved to be the most22der normal environmental conditions and after artificial exposure. Epoxy proved to be the most23effective treatment, by increasing the yarn tensile strength of 34% and the pull-out load of 138%,24followed by nano-silica (+9%; +40%). All surface treatments showed to remain effective even after25artificial environmental exposures, with a maximum reduction of yarns tensile strength of about2613%.27

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Keywords: Fabric-Reinforced Cementitious Matrix (FRCM); carbon yarn; carbon fibers; surface 29 treatments; coating; interface; bond; environmental exposure; durability. 30

1. Introduction

Fabric-Reinforced Cementitious Matrix (FRCM), also known in the international literature 33 as Textile-Reinforced Mortar (TRM), is a new class of composite material that generated 34 considerable interest as strengthening technique for concrete and masonry structures. 35 FRCMs are constituted by open grids of perpendicularly connected multifilament yarns 36 (made of carbon, glass, aramid, basalt or PBO fibers), which are applied on concrete or 37 masonry structural elements through lime or cement-based matrices [1–4]. Although the 38 use of FRCM systems as externally bonded reinforcement is nowadays a common practice 39 in civil engineering, there are still some issues that need to be addressed, such as the mod-40 est adhesion at the interface between the fabric reinforcement, made of multifilament 41 yarns, and the inorganic matrix. 42

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Copyright: © 2022 by the authors. Submitted for possible open access publication under the terms and conditions of the Creative Commons Attribution (CC BY) license (https://creativecommons.org/license s/by/4.0/). Experimental studies on FRCM systems showed that the main failure mode is by debond-43 ing and slippage of the fabric yarns within the inorganic matrix [5,6]. Slippage of multi-44 filament yarns within the matrix is emphasized in the case of yarns made up of carbon 45 fibers [7]. The absence of hydrophilic groups on the carbon chain indeed results in a rela-46 tively poor chemical compatibility between the surface of carbon fibers and inorganic ma-47 trices [8]. Moreover, the inorganic matrix (usually cement-based) is not able to fully pen-48 etrate between the filaments of the yarn, due to its high viscosity, thus allowing the inner-49 most filaments to slip over each other, showing the so-called telescopic behavior [9]. 50

Several solutions have been proposed to improve the bond at the fiber-to-matrix interface: 51 some of them directly modify the fiber surface by means of chemical-physical reactions 52 (e.g. fibers oxidation [8,10]) while others provide for partial or complete pre-impregnation 53 of the fibers with additional materials (usually organic polymers). Pre-impregnation of 54 the fibers with organic coatings has been reported by many studies as a promising tech-55 nique to improve the mechanical performance of FRCM systems [11–16]. Donnini et al. 56 found out that the use of epoxy coatings on carbon fabrics [17] or yarns [7] are very prom-57 ising in increasing the bond at the interface with the inorganic matrix, depending on the 58 level of pre-impregnation. Signorini et al. investigated the effect of epoxy resin viscosity 59 on the mechanical properties of pre-impregnated FRCM systems, showing that epoxy can 60 penetrate inside the yarn and prevents telescopic failure [18]. 61

In general, the main consequence of pre-impregnation with organic coatings is that the 62 inorganic matrix is prevented to penetrate within the filaments of the yarn and the inter-63 face bond is no longer dependent on the matrix ability to wet the single filaments but 64 rather on the interaction between the coating and the matrix. Moreover, when the single 65 filaments are embedded in an organic coating, a higher number of filaments is engaged in 66 the stress-transfer mechanisms, thus improving the yarn tensile strength and also leading 67 to a change in the failure mode (usually from fabric slippage to fabric breakage) [19]. How-68 ever, the main disadvantage of using organic coatings is the reduced ability of the com-69 posite system to withstand high temperatures [20–22]. 70

Alternatively, the use of inorganic coatings has also been investigated, in order to over-71 come the issue of polymeric matrices subject to high temperatures [23–25]. The use of in-72 73 organic coatings, usually based on silica, cement or other nano-particles [24,26,27], allows for a stronger fiber-matrix interaction, thanks to the reaction and chemical bonds that can 74 develop between the coating and the inorganic matrix, thus providing a better adhesion 75 at the fiber-to-matrix interface. The interfacial behavior between nano-silica modified car-76 bon fibers and cementitious matrices has been recently studied by Li et al. [28], showing 77 that the interfacial adhesion at fiber-matrix interface is improved compared to untreated 78 fibers, due to the formation of C-S-H gel in the vicinity of the fiber surface. The effects of 79 silica nano-coatings on carbon fibers have been also investigated by Signorini et al. [29], 80 showing a good improvement in the performance of the composite, even if lower than 81 those obtained with polymeric coatings. 82

The durability of FRCM systems is also a very topical issue [13,30–33]. Carbon fibers have 83 high resistance to chemical attacks and their properties remain almost unchanged when 84 subjected to artificial aging conditions [34,35]. However, when the fibers are coated or 85 modified with other surface treatments, the mechanical properties and durability of the 86 yarns could be compromised. 87

The objective of this study is to evaluate the effectiveness of three different surface treat-88 ments applied to multifilament carbon yarns, to improve their mechanical performance 89 and chemical-physical compatibility with cement-based matrices. The treatments investi-90 gated consist of epoxy pre-impregnation, nano-silica coating and sonication through an 91 oxidative solution. The effectiveness of the treatments was evaluated both considering the 92 tensile properties of the yarns (uniaxial tensile tests on single yarns) and the bond with a 93 cement-based mortar (pull-out tests). Moreover, mechanical properties have been also 94 evaluated after having exposed the specimens under different artificial aging conditions, 95 to verify their effectiveness even in aggressive environments (Figure 1). 96



Figure 1. Schematic representation of the experimental campaign.

2. Materials and Methods

2.1. Materials

Two different types of specimens were manufactured to evaluate the effectiveness of dif-103 ferent surface treatments on the tensile properties of single carbon yarns and on the adhe-104 sion with an inorganic cement-based mortar (as schematically reported in Figure 1). Car-105 bon yarns used in experiments have been taken from a commercially available bidirec-106 tional dry carbon fabric. The 24 K carbon yarn has a cross-sectional area equal to 1.04 mm², as reported by the manufacturer, according to ISO 527-4,5: 2021 [36]. Preparation of carbon yarns for tensile tests, test procedure and mechanical properties are reported in the 109 experimental results section. 110

Pull-out specimens have been manufactured by embedding single carbon yarns in a cube 111 of cement-based mortar (side of 40 mm), whose mix proportions and mechanical proper-112 ties have been reported in Table 1. Compressive and flexural strength of the mortar have 113 been evaluated on prismatic specimens (40 x 40 x 160 mm³) after 28 days of curing in 114 laboratory conditions (20 ± 2°C, RH=70%), according to UNI EN 1015-11 [37]. 115

Table 1. Mix proportions and mechanical properties of the inorganic matrix.

Material	CEM II/B- LL 32.5 R (kg/m ³)	CEM II/B- LL 42.5 R (kg/m ³)	CaCO ₃ 400 (kg/m ³)	CaCO3 600 (kg/m ³)	Aerial lime (kg/m³)	Water (kg/m³)	Compressive strength (MPa)	Flexural strength (MPa)
Cement- based mortar	82	165	715	205	110	260	17.95	5.66

2.2. Surface treatments

Three different surface treatments have been employed. The first one consists in the ap-119 plication of a highly flexible two-component epoxy resin (C-E), whose mechanical 120

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properties are reported in Table 2. Yarns were fully impregnated with the epoxy by means 121 of a plastic spatula, then cured at 60 °C for 24 hours. 122

The second treatment involved the application of a nano-silica coating (C-NS). Carbon 123 yarns were immersed in a nano-silica dispersion under stirring for 15 minutes and then 124 dried at room temperature. The nano-silica dispersion was obtained by sol-gel method 125 adding an acidic solution (distilled water: 65% Nitric Acid in the molar ratio 1:0.032) to a 126 98% Tetraethyl orthosilicate (TEOS) by Evonik, Ethanol (Analytical grade) solution as de-127 scribed in [38].

Finally, the third treatment (C-Ox) consists of carbon yarns sonication in a HNO₃/H₂SO₄ 129 oxidative solution for 15 minutes, followed by washing with distilled water until a decrease of the washing water pH to a value of 6. 65% nitric acid (HNO₃) and 95% sulfuric 131 acid (H₂SO₄) are purchased from Sigma-Aldrich. The oxidative solution was prepared according to [39], with 1:3 HNO₃ /H₂SO₄ volume ratio. Yarns were dried at room temperature before testing. 134

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Table 2. Physical and mechanical properties of the epoxy resin (from manufacturer).

Material	Viscosity (mPas)	Tensile strength (MPa)	Elongation at break
Elan-tech EC 98N/W52	2000-3000	0.7-0.9	60-80%

2.3. Tensile and pull-out tests

To evaluate the effectiveness of the different surface treatments on the mechanical prop-139 erties of carbon yarns and on the bond with the inorganic matrix, a total of 60 tensile and 140 80 pull-out tests have been carried out. Tensile tests on single carbon yarns were per-141 formed by using a tensile testing machine with a load bearing capacity of 50 kN, with a 142 loading rate of 0.5 mm/min, according to ISO 10406-1 [33]. FRP glass tabs were epoxy-143 bonded at the ends of the specimen to ensure a better grip during the test. A macro-exten-144 someter with a gauge length of 50 mm was positioned at the center of each specimen to 145 evaluate the elastic modulus and to measure the strain at failure (Figure 2a). Mechanical 146 parameters have been reported as the average on 5 specimens for each type. Tensile 147 strength has been calculated by dividing the tensile load by the cross-sectional area of the 148 yarn (provided by manufacturer). The elastic modulus has been calculated as the slope of 149 the stress-strain curve in the elastic branch comprised between 20% and 50% of the maxi-150 mum tensile capacity [33]. The average tensile strength (σ_{max}), ultimate strain (ε_u) and elas-151 tic modulus (E), are reported in Table 4, together with the corresponding coefficient of 152 variation (CoV). 153

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Figure 2. a) Tensile and b) pull-out test layout, c) actual pull-out test setup.

Pull-out tests were carried out on carbon yarns embedded in a cubic specimen of cementi-156 tious mortar (40 x 40 x 40 mm³). The free length is kept constant and equal to 20 mm. Pull-157 out tests were performed using a tensile testing machine with a load bearing capacity of 158 5 kN. The specimen is fixed at the bottom by a metallic frame anchored to the testing 159 machine and the upper part of the yarn is gripped and pulled in displacement control at 160 0.5 mm/min (Figure 2b). The average maximum pull-out load and the total displacement 161 corresponding to the maximum load (d_{max}) are reported in Table 5, together with the cor-162 responding coefficient of variation (CoV).

2.4. Aging Conditioning Protocol

Same tests have been carried out after subjecting the specimens to various artificial aging 165 environments, as summarized in Table 3. In the case of pull-out specimens, the artificial 166 conditioning started after 90 days of specimens curing at laboratory conditions ($20 \pm 2^{\circ}$ C, 167 RH=70%). 168

Environment	Temp	RH	Solution	Exposure Time	N° of samples (5 for each surface treatment)
None (Ref)	$20 \pm 2^{\circ}C$	70%	-	-	20 tensile tests 20 pull-out tests
Saline	$40 \pm 2^{\circ}C$	100%	2.45% NaCl + 0.41% Na2SO4	1000 h	20 tensile tests 20 pull-out tests
Alkaline	$40 \pm 2^{\circ}C$	100%	4% NaOH	1000 h	20 tensile tests 20 pull-out tests
Freeze-Thaw	-18 ± 2°C/ +40 ± 2°C	40%/ 100%	-	960 h (40 cycles)	20 pull-out tests

Table 3. Artificial aging test environments.

The first environment (Saline) comprises a 2.45% weight sodium chloride (NaCl) and 171 0.41% weight of sodium sulphate (Na₂SO₄) aqueous solution. The concentration of NaCl 172 and Na₂SO₄ was chosen according to ASTM D1141-98 [34]. In order to accelerate the aging 173 process without promoting unrealistic chemical reactions a temperature of 40 °C was cho-174 sen. 175

The second environment (Alkaline) comprises a 4% weight sodium hydroxide (NaOH) 176 aqueous solution with pH of 13. The exposure to alkaline and saline environments was 177 conducted by completely immersing the carbon yarns in the solution, while the pull-out 178 specimens were immersed for 3 cm, in order to keep the free length of the yarn out of the 179 solution (Figure 3). 180

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Freeze-thaw cycles consisted of freezing at -18 °C for more than 6 h and thawing at 40 °C 181 for about 12 hours. A total of 40 cycles have been carried out both for carbon yarns and 182 pull-out specimens. Finally, after artificial conditioning, all specimens have been dried at 183 40 °C for 24 h before testing. 184



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Figure 3. Conditioning exposure of tensile and pull-out specimens in saline and alkaline solutions 186

2.5. SEM and EDX analysis

SEM and EDX analysis have been carried out, using a FESEM ZEISS SUPRA40 with EDX-188 Detector Brucker Quantax 200-Z10, to investigate on the atomic percentages of carbon, 189 silicon, oxygen and on the surface morphology of carbon yarns after different surface 190 treatments. 191

3. Experimental results

3.1. Tensile tests

Experimental results of tensile tests on carbon yarns with different surface treatments, subjected to different environmental exposures, are reported in Table 4. 195

Table 4. Results of tensile tests on carbon yarns.

Sacimon	Environment		Tensile Strength	Variation of tensile	Elastic Modulus	Ultimate Strain
specimen	Environment		σ max,	strength	Ε	Eu
			(MPa)		(GPa)	(%)
	None (Bef)	Average	1732	-	145	1.39
	None (Ker)	CoV	6.9%		1.1%	1%
	Calina	Average	1594	-8.0%	144	1.25
C-Dry	Same	CoV	4.8%		4.7%	19.5%
	Alkaline	Average	1681	-2.9%	152	1.07
	Aikaime	CoV	10.1%		9.2%	5.8%
	None (Ref)	Average	2327	-	142	1.76%
C F		CoV	7.6%		4.5%	5.1%
	Saline	Average	2201	-5.4%	147	1.50
C-E		CoV	14.1%		5.2%	20.5%
	Allcoling	Average	2226	-4.3%	151	1.47
	Aikainie	CoV	11.4%		9.6%	17%
	None (Bef)	Average	1900	-	143	1.6
	None (Ker)	CoV	3.3%		3.4%	6.1%
CNE	Calina	Average	1672	-12.0%	149	1.08
C-INS	Same	CoV	10.4%		3.4%	9.9%
	Allcoling	Average	1651	-13.1%	150	1.24
	Aikainie	CoV	3.3%		4.6%	12.3%
	None (Def)	Average	1667	-	139	1.39
C-Ox	mone (Ker)	CoV	8.4%		4.3%	12%
	Saline	Average	1626	-2.4%	155	1.10



Figure 4. Tensile strength of carbon yarns with different surface treatments exposed to different environments.

Carbon yarns impregnated with epoxy resin (C-E) showed the highest tensile strength, 201 equal to 2327 MPa, which corresponds to an increase of the tensile strength of about 34%, 202 if compared to untreated varns. The impregnation with nano-silica dispersion also leads 203 to a slight increase of the tensile strength, of about 10%, while apparently the oxidative 204 treatment has barely influenced the mechanical properties of the yarn. The ability of 205 organic and inorganic coatings to improve the mechanical properties of multi-filament 206 yarns has been reported in different studies, and it is attributed to the stress transfer 207 increase between single filaments [35,36]. The effectiveness of a coating to improve the 208 yarn tensile strength depends on its ability to simultaneously engage the single filaments 209 of the yarn during the test. This is in accordance with the results obtained by C-E Ref and 210 C-NS Ref. Different failure modes can also be observed for treated carbon yarns. C-211 Dry_Ref, C-NS_Ref and C-Ox_Ref yarns failed before all the carbon filaments have 212 reached their maximum tensile strength (Figures 5a,c,d), suggesting that the nano-silica 213 coating is not able to effectively activate all the yarn's filaments during tensile test. On the 214 contrary, C-E_Ref yarns showed an abrupt and simultaneous breakage of all the yarns' 215 filaments (Figure 5b). These results confirm the superior ability of the epoxy resin to 216 uniformly distribute the stress between the single filaments, thus increasing the yarn 217 tensile strength. The elastic modulus seems not be particularly affected by the surface 218 treatment employed. 219



Figure 5. Failure modes: a) C-Dry_Ref, b) C-E-Ref, c) C-NS_Ref, d) C-Ox_Ref.

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Looking at the results of durability tests (Figure 4), it can be observed that the tensile 222 strength of carbon yarns is slightly affected by the exposure to saline and alkaline 223 environments, regardless of the type of surface treatment applied. The most significant 224 reduction of tensile strength is observed for C-NS yarns. After immersion in the alkaline 225 solution, the tensile strength of C-NS yarn showed a decrease of about 13%. Similar results 226 are obtained after immersion in the saline solution, with a 12% decrease of the tensile 227 strength. The nanosilica coating is therefore effective in increasing the yarn tensile 228 performance in laboratory environmental conditions (C-NS) but it seems to suffer the 229 exposure to saline and alkaline solutions. This effect was also found in other studies [42], 230 where the ineffectiveness of nanosilica coatings is attributed not only to the reduced 231 particle size (50 nm) but also to to a partial wash out of the small particles when the 232 reinforcement is immersed in the fresh cementitious paste washing. In this study, 233 immersion of carbon yarns in saline and alkaline solution at 40 °C could have caused this 234 washout phenomenon. However, since SEM analysis of the yarn surface after exposure to 235 various artificial environments have not yet been performed, these results will have to be 236 confirmed by more detailed investigations. 237

3.2. Pull-out tests

Results of pull-out tests for all carbon yarns tested are reported in Table 5, while loaddisplacement curves are reported in Figure 6. 240

Specimen	Environment		Max pull- out load (N)	Variation of max load (%)	Displacement at max load (mm)
		Average	321	-	0.86
	None (Ref)	CoV	4%		13%
	Callin a	Average	329	+2.5	0.87
	Saline	CoV	11%		11%
C-Dry	A 11 - 11	Average	307	-4.4	0.69
	Alkaline	CoV	2%		23%
	Encore There	Average	290	-9.7	0.96
	Freeze-Thaw	CoV	23%		16%
	None (Bef)	Average	766	-	4.59
	None (Ker)	CoV	2%		5%
	Calina	Average	823	+7.4	4.98
C-E	Same	CoV	2%		17%
	A 11. a 1i. a a	Average	857	+11.9	4.99
	Alkaline	CoV	2%		12%
	Encore Theory	Average	660	-13.8	4.23
	Freeze-Thaw	CoV	5%		6%
	Name (Def)	Average	452	-	0.94
	None (Ker)	CoV	2%		7%
	Calina	Average	397	-12.2	0.7
CNC	Same	CoV	8%		3%
C-IN5	Allealing	Average	436	-3.5	0.71
	Аткаппе	CoV	10%		19%
	Eroozo Thou	Average	395	-12.6	0.91
	Fleeze-Illaw	CoV	6%		11%
	None (Ref)	Average	413	-	0.94
$C \cap \mathbf{v}$	inone (iter)	CoV	5%		7%
C-Ux	Salina	Average	364	-11.9	0.94
	Jaime	CoV	10%		5%

Table 5. Results of pull-out tests of carbon yarns subjected to different aging protocols.

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	Allcoling	Average	407	-1.5	0.78
Aikainie		CoV	5%		7%
	Encore Theory	Average	301	-27.1	0.74
	Freeze-Thaw	CoV	8%		5%

At first, it can be observed that all the surface treatments investigated in this study are able to increase the maximum pull-out load with respect to reference yarns.

Dry carbon yarns fail at low load values, due to poor chemical-physical interaction 244 between dry carbon filaments and the cementitious matrix. The failure is due to the 245 breakage of some external filaments of the yarn and consequent slippage of the inner ones 246 (telescopic effect). This failure mode can be observed in the broken specimen of Figure 7a, 247 where only few external filaments remained attached to the inorganic matrix after pullout test. 249

The use of epoxy-based coating showed to be the most effective treatment, by increasing 250 the pull-out load of about 138% with respect to dry carbon yarns. This is due to the greater 251 and more homogeneous stress distribution between all the filaments of the yarn (as also 252 observed in tensile tests), and to the high friction which develops at the epoxy-to-253 inorganic matrix interface. These results are in agreement with some findings from the 254 literature, where an average 2-3 times increase of the mechanical performance of FRCM 255 composites with polymer impregnated carbon yarns, compared to dry fiber yarns, is 256 observed [7,43]. Similar results have been found by Signorini et al. for epoxy coated glass 257 fibers [13]. 258

The maximum pull-out load of C-E samples occurred for very large displacements. In fact,259once the yarn detaches from the inorganic matrix (first peak in Figure 6, C-E), it starts to260slip within the matrix and the pull-out load increases up to reach the maximum value.261This stress-hardening behavior can be explained by looking at the surface of the C-E yarns262after pull-out (Figure 7b), which shows that the epoxy coating has been partially removed263by friction with the inorganic matrix.264

The maximum pull-out load of carbon yarns treated with oxidative solution (C-Ox) and 265 nano-silica dispersion (C-NS) was respectively 28% and 40% higher than that of dry yarns 266 (C-Dry). Looking at the load-displacement curves, C-Dry, C-NS and C-Ox specimens 267 showed a similar pull-out behavior, characterized by a first linear increase of the load, 268 followed by a quick load decrease after the peak. Since neither the nano-silica coating nor 269 the oxidation treatment influenced the graph shape in the post-debonding region, it can 270 be assumed that these treatments did not affect the frictional shear stress at the composite 271 interface. Therefore, the higher peak loads obtained in both cases are the consequence of 272 a higher chemical bond with the cementitious matrix. For C-NS yarns this can be 273 attributed to the chemical reaction of silica particles with the Ca(OH)2 of the cementitious 274 mortar, forming a calcium silicate hydrate (C-SH) layer in the proximity of the fibers [26]. 275 For what it concerns C-Ox samples, the oxidation process is expected to modify the carbon 276 fibers surface, with the formation of oxygen-containing functional groups which help the 277 wettability of the fibers by the cementitious mortar [44]. Some studies in the literature 278 show the effectiveness of different oxidative treatments in improving the bond strength 279 between carbon fibers and cementitious mortars, and therefore also the mechanical 280 properties of the composite. However, it is difficult to compare the results of this study to 281 others from the literature, due to a large number of variables. Some studies indeed refer 282 only to short carbon fibers (instead of multifilament yarns) [44,45] while others use 283 different oxidation processes or different setups for mechancal tests [46,47]. 284

Regarding the oxidative treatment used in this study, some observations on its 285 effectiveness are reported in Section 3.3, following SEM and EDX analysis on the yarn 286 surface. 287

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Figure 6. Load-displacement curves of pull-out tests.



Figure 7. a) Contact surface between dry carbon yarns (C-Dry) and inorganic matrix after pull-out tests, b) external surface of epoxy coated carbon yarn (C-E) after pull-out, c)matrix breakage.

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Regarding the results of pull-out tests after exposure to various artificial environments, a300graphical representation of the outcomes is reported in Figure 8. It can be observed that,301regardless of the environmental exposure, the unmodified yarns always show the lowest302pull-out load. Moreover, the different environments do not seem to significantly affect the303results of pull-out tests, regardless of the type of surface treatment employed.304

It is interesting to note that C-E yarns subjected to alkaline and saline environments show 305 a slight increase of the pull-out load, suggesting that the epoxy resin is able to further 306 protect the carbon fibers from aggressive environments and that immersion in solution at 307 40°C has even increased the bond at the epoxy-matrix interface. However, in some cases 308 a brittle failure at the yarn-matrix interface was observed, with the complete separation of 309 the matrix into two parts (Figure 7c). This phenomenon can also be observed in some load-310 displacement curves, with an abrupt decay of the load corresponding to the matrix 311 breakage (Figure 6, C-E_Sal, C-E_FT). This is a confirmation that the superior properties 312 of epoxy-coated carbon yarns are mainly due to the friction that develops during the 313 slippage of the yarn within the matrix more than to the chemical adhesion between the 314 cured epoxy resin and the matrix (very low). Therefore, the presence of an epoxy coating 315 can act as separating layer and promote delamination failures, as also observed in other 316 studies [13,48]. 317

Surface treatments based on nano-silica (C-NS) and fibers oxidation (C-Ox) showed to be 318 adequately resistant to the aggressive environments, confirming their superior properties 319 compared to dry yarns. In this study the most degrading environment was that of freeze-320 thaw cycles, which caused a decay of the pull-out load for all the specimens investigated 321 (up to -27% for C-Ox specimens). This can be attributed to internal damage of the 322 cementitious matrix, as well as to an incomplete curing of the matrix due to the low 323 temperature and humidity of the conditioning environment, which could have led to 324 premature failure (although the matrix did not show any significant cracks due to freeze-325 thaw cycles). 326



Figure 8. Pull-out load of carbon yarns with different surface treatments after artificial exposure.

3.3. EDX and SEM analysis

The results of EDX analyses on the surfaces of C-Dry, C-NS and C-Ox yarns are reported331in Table 6. The atomic percentage of carbon ranges from 83% up to 94%. The presence of332oxygen atoms can be attributed to the organic sizing applied to carbon filaments during333the manufacturing process of the fabric.334

It is interesting to observe that EDX analyses conducted on C-Ox yarns excluded the formation of new oxygen bearing groups on the carbon fiber surface (Table 6). A possible explanation of the improved interaction observed for C-Ox samples in pull-out tests is 337

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that the chemical treatment was not sufficient to promote the oxidation of the carbon back-338 bone, but it was able to attack the fibers surface causing an improvement of its roughness. 339 An experimental study conducted on cement-based composites reinforced with carbon 340 fibers (although short fibers) treated with concentrated acid, attributed the improved 341 interaction between the matrix and the fibers to the formation of hydroxyl and carboxyl 342 groups on the fibers surface [45]. However, in this study the results of SEM analysis 343 showed no significant difference on the surface morphology of the fibers before and after 344 the treatment with the oxidative solution (Figures 9a,b). A further explanation is that no 345 oxidation took place, but the acid was able to catalyze other chemical reactions which did 346 not change the chemical composition of the fibers but could modify the oxygen bearing 347 groups on the fibers surface. Acids are known to catalyze different chemical reactions, 348 however, since the exact chemical composition of the sizing is unknown, it is not possible 349 to state which phenomenon may have occurred. Moreover, because no clear FTIR 350 spectrum of the fibers before and after chemical treatment could be acquired this 351 hypothesis cannot be confirmed. 352

On the other hand, the clear increase in the percentage of oxygen and silicon atoms 353 observed for C-NS samples, compared to the untreated yarn, confirms the presence of 354 nano-silica particles between the yarn filaments. However, SEM analysis (Figure 10) 355 shows that the nano-silica coating is not uniformly distributed between the filaments, thus 356 forming a discontinuous layer on the fibers surface. This is probably due to the manual 357 impregnation process, which is not able to adequately control the uniformity of the 358 application. 359

Sample	C [At %]	Si [At %]	O [At %]
C-Dry	93.47	0.23	6.31
C-NS	83.9	1.47	14.63
C-Ox	94.11	0.12	5.76

 Table 6. Atomic percentages of carbon, silicon and oxygen detected by EDX analyses on the



surfaces of the yarns C-Dry, C-NS and C-Ox

Figure 9. SEM images of a) C-Dry, and b) C-Ox multifilament yarns. Magnification 5.00 Kx.

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Figure 10. SEM images of C-NS yarns. Magnification (a) 2.00 Kx and (b) 5.00 Kx.

4. Conclusions

Based on the results of this experimental investigation, the following conclusions can be drawn:

- Pre-impregnation of multifilament carbon yarns with epoxy proved to be the 370 most effective treatment, capable of increasing both the yarn tensile strength and 371 the bond with the cement-based mortar. Epoxy was able to increase the carbon 372 yarn tensile strength of about 34% and the pull-out load of about 138%. These 373 effects, as well known from the literature, can be attributed to the ability of the 374 low-viscosity epoxy to penetrate between single filaments of the yarn, thus guar-375 anteeing a more homogeneous stress distribution through the yarn cross-section. 376 However, it must be remembered that the use of organic polymers remains a 377 weakness as regards the mechanical behavior of the composite when exposed to 378 high temperatures. 379
- Nano-silica coating was less effective than epoxy, but still able to increase the yarn 380 tensile strength of 10% and the pull-out load of about 40%. However, the effectiveness of this treatment can be improved by optimizing the manufacturing process, trying to ensure a more homogeneous distribution of the particles on the 383 yarn surface. 384
- The oxidation of carbon fibers with HNO₃/H₂SO₄ solution seems not to substantially modify the mechanical properties of the carbon yarns. SEM analyses did not show significant changes in the surface of the carbon filaments after the oxidation process. However, this treatment was able to increase the pull-out load of about 28%. Further analyses are certainly needed to better investigate this aspect.
- Artificial aging in saline and alkaline environments caused only a slight reduction 390 of the yarns tensile strength, which was always lower than 13%, regardless of the 391 type of surface treatment applied. 392
- Pull-out tests carried out after exposure of the specimens in saline and alkaline 393 environments showed no significant decrease of mechanical performances. Also 394 in this case, carbon yarns with epoxy impregnation showed the highest load val-395 ues. Exposure to freeze-thaw cycles caused the greatest reduction in the pull-out 396 load (between -10% and -27%), probably due to internal damage of the inorganic 397 matrix (which in some cases broke in half during the test), rather than to deterio-398 ration of the carbon yarn.

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