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# Effect of high temperature exposure on epoxy-coated glass textile reinforced mortar (GTRM) composites

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#### Abstract

An experimental investigation on the mechanical performance of epoxy-coated Alkali-Resistant (AR) glass textile reinforced mortar subjected to elevated temperature is presented. Two epoxy coatings are considered, which differ by the hardening agent alone. After 56 days dry curing, specimens are heated up to four different temperatures. After cooling down to ambient temperature, specimens are assessed in uni-axial tensile test according to Annex A of AC434. First cracking strength and elongation, ultimate tensile strength and elongation, cracked and uncracked moduli, transition point location and energy dissipation capability are evaluated. It is found that, in the explored temperature range, degradation is surprisingly mild and strongly dependent on the resin which is taken as coating agent. Indeed, temperature exposure may lead to strength enhancement. This positive outcome takes place at the expense of ductility and it is traced back, through Differential Scanning Calorimetry (DSC), to a postcuring process. Nonetheless, energy dissipation still decreases with temperature and, remarkably, with the same power-law behaviour for both resins. Such behaviour is compatible with a cumulative Weibull distribution, that is adopted in thermal damage models for resins, and it indicates that the underlying damage mechanism indeed operates on the resin at the fabric-to-matrix interface.

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#### 1. Introduction

The possibility of high temperature exposure poses a serious limitation to the applicability of organic matrix reinforcing systems, such as fibre-reinforced polymers (FRPs). Indeed, exposing FRP systems to temperatures in excess of or even close to the glass transition temperature  $T_q$  produces a substantial and sudden drop in the mechanical response [10, 6]. This behaviour, that rapidly leads to delamination and failure, is all the more undesired in consideration of the fairly low transition temperature  $T_g \approx 80$  °C characterizing most organic resins. In this respect, Textile Reinforced Mortar/Cement (TRM/TRC) and Fabric Reinforced Cementitious Matrix (FRCM) composite materials exhibit vastly superior thermal stability, in light of the adoption of an inorganic matrix. On the other hand, the bond strength between the fabric and the inorganic 11 matrix is generally weak and this leads to poor mechanical performance and 12 a generally inconsistent failure pattern [2, 21]. Improvement in the matrix-to-13 fabric bond may be obtained by adopting inorganic [13, 31] or organic [29, 9, 19] coatings. Consequently, investigation of the effect of temperature exposure on TRM is complicated by the need to consider the whole composite package, which 16 consists of the matrix, the fabric and the coating. 17 A large body of literature is devoted to the characterization of FRP systems 18 subjected to elevated temperature, see, for instance, [10, 7, 4, 28, 16, 15] and references therein. Conversely, a limited number of studies is available concern-

subjected to elevated temperature, see, for instance, [10, 7, 4, 28, 16, 15] and references therein. Conversely, a limited number of studies is available concerning the effect of high temperature exposure on TRM and FRCM and these are mainly focused on carbon and PBO fabrics [24, 34]. In this framework, a crucial issue that requires careful investigation is the role played by high temperature exposure on the adhesive behaviour of the laminates at the mortar-to-substrate interphase, as discussed by Ombres [23] and Maroudas and Papanicolaou [18] for concrete and masonry structures, respectively. The former study presents single-lap shear tests on PBO-FRCM laminates applied on concrete supports

and reports that conditioning at 50 °C and 100 °C affects not only the loadbearing capacity of the composite, which decreases from 25 to 40% depending on the number of layers, but also the failure and the delamination modes. Indeed, failure statistically changes from matrix-to-fabric slippage for the unconditioned samples to debonding at matrix-to-concrete interphase for the conditioned ones. 32 Analogously, in the contribution by Maroudas and Papanicolaou [18], single-lap 33 shear tests are conducted on G-TRM thin laminates applied to brick panels and exposed to temperatures up to 300 °C. It is shown that, if the temperature exceeds 100 °C, failure is mainly triggered by adhesive debonding at the masonry-to-mortar interphase and strongly affected by the deterioration of the 37 ultimate strength of the bare glass fabric. Furthermore, Raoof and Bournas [26] 38 and Bisby et al. [3] assess the response in bending of TRM and FRP reinforcing systems subjected to high temperatures, while Tetta and Bournas [32] considers jacketing. Trapko [33] compares FRP and FRCM confined concrete elements exposed to temperatures up to 80 °C for 24 h. Already at 40 °C compressive strength of FRP jacketed elements is reduced by 20% and at 80°C ductility 43 drops by 50%, as opposed to a 11% loss encountered for FRCM. de Andrade et al. [8] investigate double-sided pull-out strength of an epoxy coated carbon yarn after 120 min exposure at 100 °C, 150 °C, 200 °C, 400 °C and 600 °C. Maximum pull-out force and pull-out work are computed and compared with the dry yarn. Interestingly, an increase in mechanical performance of the coated 48 specimens is observed after heating at temperatures up to 150 °C, that is ascribed to a "polymer interlocking mechanism in the yarn-matrix interface, which is generated during the heating and cooling of the polymer yarn coating". In 51 Rambo et al. [25], uni-axial tensile tests of basalt textile reinforced plates are 52 conducted. The basalt fabric is coated with styrene-acrylic latex and refractory 53 concrete is adopted as matrix. Plates are exposed for 60 min at temperatures in the range 75–1000 °C. It is found that performance loss is mild up to 200 °C and it is concluded that "the presence and the type of coating can become a de-56 terministic factor in the tensile response of the composite submitted to elevated temperatures". Recently, Donnini et al. [9] present experimental and numeri-

cal results on the mechanical performance of dry and epoxy-plus-sand coated carbon FRCM composites under uni-axial tension and double-shear bond test. Beside ambient temperature, 120 min exposure at 80 °C and at 120 °C are considered. It is worth emphasizing that mechanical tests are carried out inside the climatic chamber, where specimens are exposed at high temperature. An impressive 70% drop in the ultimate tensile strength (and a 54% elongation loss) is determined with respect to the ambient condition already at 80 °C. In general, the existing literature lacks from detailing the thermo-physical properties of the coating agent, that is usually adopted taking an out-of-the-67 box approach. In this work, we focus on the role of epoxy coating on the thermal deterioration of the matrix-to-fabric interface. To this aim, the same 69 pair of epoxy resins considered in [19] are adopted, which differ only by the hardening agent. This feature, however, leads to important differences in terms of thermo-physical properties that exert a profound influence on the behaviour upon high temperature exposure [12]. To avoid degradation of the lime mortar and restrict attention to the epoxy coating, temperature exposure is limited in 74 the range 20–250 °C and 56-day dry curing is adopted. Mechanical performance is assessed in uni-axial traction of rectangular coupons according to Annex A of the guidelines [14]. DSC analysis supports the conclusion that, depending 77 on the resin formulation, high temperature exposure may benefit strength (at

# 2. Materials and methods

83 2.1. Materials

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- 84 2.1.1. Reinforcing fabric and inorganic matrix
- The commercially available Alkali Resistant Glass (ARG) fabric Zirconglass
  Wire<sup>©</sup> RV320-AR (Fibre Net Spa) is adopted as fabric reinforcement. This is
  a balanced bi-axial open-squared mesh whose 19% weight content of Zirconium
  Oxide (ZrO<sub>2</sub>) imparts resistance to the alkaline mortar environment. The main

the expense of ductility), inasmuch as it promotes cross-linking in a post-curing

process. This mechanism is likely to explain the outcome of the double pull-out tests carried out in [8] and of the uni-axial traction of plates described in [25].

Table 1: ARG Fabric mechanical properties (tex = g/km)

Characteristic	Unit	Value
Yarn count	tex	1200
Net specific weight per unit fabric area	$g/mm^2$	300
Fabric specific weight	$\rm g/cm^3$	2.50
Grid spacing (square grid)	mm	12
Equivalent thickness, $t_f$	mm	0.06
Ultimate strength along warp (with epoxy)	MPa	1200
Ultimate elongation along warp	mstrain	20
Elastic modulus	GPa	74

Table 2: Mortar properties

Characteristic	$\mathbf{Unit}$	Value
Nominal setting water content	%	21.2
Final density	$\rm g/cm^3$	1.58
Min. compression strength after 28 days	MPa	15.0
Min. flexural strength after 28 days (EN 196/1)	MPa	5.0
Min. support adhesion strength after 28 days	MPa	1.0
Aggregate maximum size	mm	1.4
Compression elastic modulus (EN 13412)	GPa	9.0

- properties of the fabric, as given by the manufacturer, are collected in Table 1.
- A pre-mixed natural hydraulic lime (NHL) mortar GeoCalce Fino<sup>©</sup> (Kerakoll
- <sup>91</sup> SpA), aimed at structural purposes, constitutes the inorganic embedding matrix.
- Table 2 gathers the main properties of this fine-grained repair mortar as given
- by the manufacturer.
- 94 2.1.2. Fabric sizing and coating
- ARG fabric is subjected, as received, to a preliminary sizing treatment to en-
- hance chemical compatibility with the epoxy coating. Following the procedure

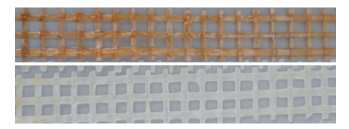


Figure 1: Coated fabric before the heat treatment (upper: ER, lower: EW)

Table 3: Hardening agents datasheet

Characteristic	Unit	m-PDA	DETA
Physical form	-	Pellets	Liquid
Formula	-	$C_6H_8N_2$	$C_4H_{13}N_3$
Melting point	$^{\circ}\mathrm{C}$	$63 \div 65$	-40
Flash point	$^{\circ}\mathrm{C}$	175	94
Boiling point	$^{\circ}\mathrm{C}$	$282 \div 284$	$200 \div 204$

described in [19], fabric is functionalized by immersion in a 2% vol. aqueous solution of (3-Aminopropyl)triethoxysilane (99 %, Sigma-Aldrich), which takes on the role of coupling agent. Care is taken to avoid organic solvents which may damage the thermoplastic stitches that hold the unwoven fabric. Fabric is then 100 dried in ambient air. Fabric epoxy coating is obtained from high-purity bisphe-101 nol A diglycidylether resin D.E.R. 332, (DOW Chemicals, hereafter "DER"). 102 Two coatings are considered, named ER and EW, which only differ by the curing 103 agent: ER exploits the aromatic hardener m-phenylenediamine (99%, Acros Organics hereafter "m-PDA"), while EW adopts the aliphatic diethylenetriamine 105 (99%, Alfa-Aesar hereafter "DETA"). Table 3 presents the main characteristics 106 of the curing agents as declared by the producers, with particular emphasis on 107 thermal properties. The coated fabric is laid on a polypropylene sheet to pre-108 vent warping, which may hinder the lamination process, and then it is allowed to set for 7 days at laboratory conditions (Fig.1). 110

# 2.1.3. Specimen manufacturing

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1-ply ARG-TRM coupons are manufactured on an individual basis by means 112 of a dismountable polyethylene formwork, following a well-established and reli-113 able manufacturing protocol, see [30, 22]. The lubricated surface of the formwork 114 is segmented by 3 mm-thick laths, equally spaced according to the specimen 115 width. In between adjacent laths, uniformly-thick mortar layers are laid out. Indeed, the top surface of the laths provides an easy reference for scraping off 117 the mortar in excess of 3 mm. Cut-to-size glass fabric is laid on top of the fresh 118 mortar and gently pressed on it. Then, a second array of polyethylene laths is 119 pinned on top of the first to provide reliable fabric placing as well as reference



Figure 2: Manufacturing process of the specimens: application of the second set of constraining laths for uniform placement of the second layer of mortar

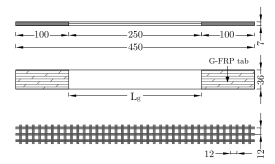


Figure 3: Coupon and fabric mesh geometries

for laying out the second mortar layer (Figure 2). At the final stage of their placing, laths are covered with paper adhesive tape to ease specimen stripping. 122 A minimum of four specimens is considered for each test group. 7-day moist 123 curing is followed by dry curing at room temperature for 56 days in total. In-124 deed, curing time is proven to deeply affect the mechanical performance of lime and cement-based composites [22] and their resistance to aggressive environ-126 ments [20]. The specimen geometry (coupon) is schematically drawn in Fig.3 127 alongside the fabric mesh size. After heating and natural cooling, a pair of ex-128 ternally bonded 100-mm G-FRP tabs is glued at both ends of each specimen to 129 accommodate the clamps of the testing machine.

Table 4: High temperature exposure conditions in the literature

Ref.	Temp. [°C]	Exp. time [min]	Composite	Test
Xu et al. [34]	120, 200	30, 90	CFRCM+epoxy	3-point bending
Donnini et al. [9]	20, 80, 120	100	CFRCM+epoxy	uni-axial traction, double shear bond
de Andrade et al. [8]	100, 150, 200, 400, 600	120	CFRCM+epoxy	double sided pull- out
Trapko [33]	40, 60, 80	1440	CFRCM	compression of confined cylinders
Rambo et al. [25]	75, 150, 200, 300, 400, 600, 1000	60	basalt+latex+FRCM	traction of plates
Ombres [23]	20, 50, 100	480	PBO-FRCM	single-lap shear
Ombres [24]	20, 50, 100, 150, 200, 250	1200	PBO-FRCM	compression on confined cylinders
Maroudas and Papani- colaou [18]	20, 100, 200, 300	1200	GFRCM	single-lap shear

# 2.2. High temperature exposure

After curing, coupons undergo a heating treatment in a Binder WTC oven. 132 A 4°C/min heating ramp is applied until either of four different target temperatures is reached, namely 100, 150, 200 or 250 °C. The set of target temperatures 134 is chosen to induce coating degradation only. Indeed, according to [5], fabric 135 composites in a cement-based matrix perform well up to 450 °C. Once the target 136 temperature is attained, isothermal conditions are maintained for 120 min. It 137 should be observed that heating time and target temperature are not standard and indeed they vary greatly across the relevant literature, as summarized in 139 Tab.4. Specimens are then moved to room temperature  $(20 \pm 2^{\circ}C)$  and left to 140 cool down in a natural cooling process, as in [8]. 141

#### 3. Experimental characterization

#### 3.1. Optical investigation

Preliminary visual investigation of the specimens after heating is illustrated in Fig.4. In particular, both epoxy coatings, when exposed to temperatures

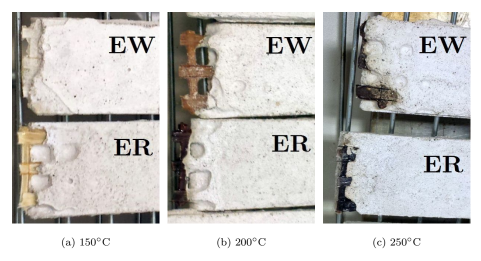


Figure 4: Mortar and fabric after temperature exposure: it clearly appears that both epoxy coatings oxidise above 150  $^{\circ}\mathrm{C}$ 

higher than 150 °C, appear oxidized and blackened (compare with the uncoated fabric shown in Fig.1), while little to no effect is visible at lower temperature.

Mortar appears unaffected by any temperature.

# 3.2. Differential Scanning Calorimetry

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A Differential Scanning Calorimetry (DSC) analysis (TA DSC 2010, TA Instruments, New Castle, DE, USA) is performed on both epoxy resins, EW and ER, in a single heating ramp, starting from 0 °C up to 250 °C, with a heating rate of 10 °C/min, under nitrogen flow. The analysis is conducted at two different stages, namely immediately after resin preparation ("as mixed" condition) and after two-week curing at ambient temperature. Comparing the heating enthalpy developed in the two conditions yields the *conversion degree*, that measures the extent to which cross-linking may occur at ambient temperature.

#### 3.3. Uni-axial monotonic tensile test

Following the guidelines [14], mechanical performance is assessed in uniaxial tensile test. A Instron 5567 electromechanical Universal Testing Machine (UTM) is employed. The UTM is equipped with a 30 kN load cell and a pair of wedge clamps which, as specified in [14, §A2.2], "shall apply sufficient lateral

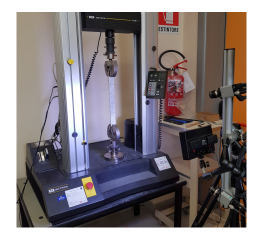


Figure 5: Uni-axial tensile test set-up with DIC monitoring of the speckled specimen

pressure to prevent slippage between the grip face and the coupon". Besides, rotationally self-aligning grips are adopted, "to minimize bending stresses in the coupon". Tests are performed under displacement control at a nominal displacement rate of 0.5 mm/min, that complies with the elongation rate proposed by the RILEM committee [27].

As already pointed out in [20], for the correct determination of elastic moduli, transition points and of the strain evolution during testing, the sliding displacement occurring in the wedge clamps needs to be subtracted from the nominal elongation ramp. To this aim, a Dantec Dynamics Q400 Digital Image Correlation (DIC) system is employed to measure the actual specimen displacement. Indeed, comparison of the nominal data with the DIC-measured elongation reveals a  $8 \div 10\%$  discrepancy in strain evaluation. The test set-up is shown in Fig.5.

#### 176 4. Results

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# 4.1. Mechanical performance

Fig.6 presents the mean strength curve for all test groups. As customary, strength is reported to the coated fabric cross-section and strain is normalized against the gauge length  $L_g$ . It immediately appears that the EW group performance is significantly impaired by the heating conditioning at any temperature,

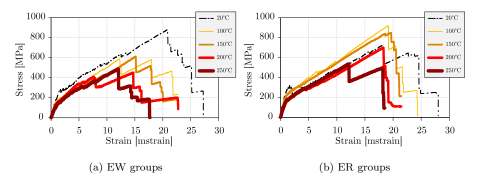


Figure 6: Mean stress-strain curve for the control (black, dashed-dotted line) and the exposed groups (solid lines with increasing thickness in dependence of the temperature exposure), namely  $100\,^{\circ}\text{C}$  (yellow),  $150\,^{\circ}\text{C}$  (orange),  $200\,^{\circ}\text{C}$  (red) and  $250\,^{\circ}\text{C}$  (amaranth)

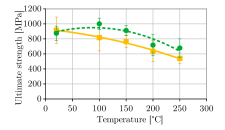


Figure 7: Mean ultimate tensile strength as a function of the exposure temperature for ER (circles, green) and EW (squares, yellow).  $\pm 1$  standard deviation bars and parabolic curve-fits are also presented

Table 5: Mean first cracking strength (FCS), ultimate tensile strength (UTS), uncracked and cracked moduli for the EW group as a function of the exposure temperature. CV is the coefficient of variation

${f T}$	FC	$\mathbf{S}$	UT	'S	$E_t^*$		$ E_{j} $	
[°C]	$\mu(f_{cr})$	CV	$\mu(f_u)$	CV	$\mu(E_f^*)$	CV	$\mu(E_f)$	CV
[ 0]	[MPa]	[%]	[MPa]	[%]	[GPa]	[%]	[GPa]	[%]
20	271.0	23.1	915.6	19.2	249.7	20.5	30.8	23.1
100	163.1	14.0	816.3	21.6	190.9	12.6	37.4	14.0
150	154.2	15.0	762.7	10.7	193.8	38.1	34.7	1.5
200	165.1	42.0	634.8	22.0	162.8	21.7	39.0	42.0
250	168.8	51.2	540.7	13.4	160.8	41.0	29.8	51.8

Table 6: Mean first cracking strength (FCS), ultimate tensile strength (UTS), uncracked and cracked moduli for the ER group as a function of the exposure temperature. CV is the coefficient of variation

$\overline{\mathbf{T}}$	FC	S	UT	$\overline{\mathbf{S}}$	$ E'_{i} $	k f	$E_j$	ŗ
[°C]	$\begin{array}{ c c } \mu(f_{cr}) \\ [\text{MPa}] \end{array}$	CV [%]	$\mu(f_u)$ [MPa]	CV [%]	$\begin{array}{ c c } \mu(E_f^*) \\ \text{[GPa]} \end{array}$	CV [%]	$\mu(E_f)$ [GPa]	CV [%]
20	192.4	24.4	875.0	10.9	319.7	21.1	31.0	15.0
100	183.3	16.4	1000.3	7.6	286.1	18.6	40.1	7.2
150	225.5	5.9	912.0	7.1	250.9	10.1	35.3	10.2
200	135.4	30.3	717.6	19.3	156.4	1.8	32.4	22.2
250	124.5	29.5	678.0	18.1	187.4	18.7	37.2	6.8

while the ER groups exhibit a mixed response. Results in terms of first crack-182 ing strength, ultimate tensile strength (UTS), cracked and uncracked moduli 183 are summarized in Tab.5 for EW and in Tab.6 for ER. This behaviour is better illustrated by the curves of Fig.7, which compare the mean ultimate tensile 185 strength (UTS) across the two groups. Indeed, while the mean UTS across the 186 EW group decreases monotonically with the exposure temperature, it increases 187 significantly in the ER-100 group and marginally in the ER-150 group, before 188 it starts to decay. Data scattering for elastic moduli is presented in Fig.8 as 189 a function of the conditioning temperature. Parabolic curve-fitting shows that 190 scattering decreases upon temperature exposure up to a critical temperature 191 that is connected to a post-curing phenomenon, as discussed in Sect.4.2. 192

In general, even for EW, heat conditioning has a surprisingly limited effect on the performance decay of the coated fabric, especially when results are compared

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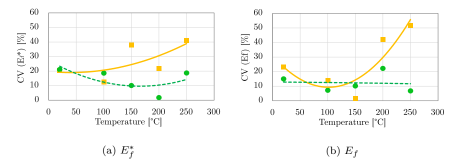


Figure 8: Coefficient of Variance (CV) for the uncracked (a) and cracked (b) secant moduli as a function of the exposure temperature for EW (orange) and ER (green) alongside its parabolic curve-fit. It is seen that post-curing positively affects data scattering as well as absolute performance

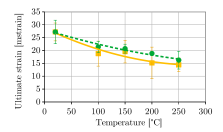


Figure 9: Mean ultimate strain values as a function of the exposure temperature for ER (circles, green) and EW (squares, yellow).  $\pm 1$  standard deviation bars and parabolic curve-fits are also presented

with the existing literature. In fact, although the performance pattern of ER is similar to that observed in [8, Fig.5] in the context of a double-sided pull-out test of a epoxy coated carbon multi-filament yarn, it should be remarked that, in the absence of a DSC analysis, the polymer coating adopted there seems exceptionally thermostable, for it cross-links at 160 °C and "the polymer film remained stable at temperatures up to 200 °C".

Fig.9 presents a similar comparison of the mean ultimate strain at failure and it shows that ductility decreases with temperature through a similar trend for both coatings.

#### 4.2. Thermal analysis

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The DSC analysis reveals an exothermic peak for both resins, associated to two-week post-curing in ambient conditions. In order to estimate the con-

Table 7: Specific enthalpy associated to curing  $(H_{curing})$  of as-mixed and two-week-cured EW and ER resins and corresponding conversion degree.

Resin	$H_{ci}$	uring [J/g]	Conversion degree
itesiii	as mixed	two-week cured	[%]
$\overline{\mathrm{EW}}$	387	15	96
$\operatorname{ER}$	379	123	67

version degree that could be achieved, the specific enthalpy measured from the DSC thermograms of Fig.10 in the two-week-cured group is compared to the 208 corresponding value obtained in the "as-mixed" group, as summarized in Table 209 7. For EW, the specific enthalpy associated to curing is located at 387 J/g in the 210 as-mixed condition and plunges to  $15 \,\mathrm{J/g}$  (corresponding to less than 4%) after 211 two-week curing at ambient temperature. Consequently, two-week curing lends 212 a conversion degree of about 96% when DETA is employed as curing agent. The same procedure applied to ER (that is when m-PDA acts as curing agent) 214 reveals that the conversion degree achieved after two week curing at ambient 215 temperature is much lower: about 67%. In fact, aliphatic amines allow curing at 216 room temperature, whereas aromatic amines usually require a high-temperature 217 treatment to achieve full conversion. However, aromatic amine-cured systems can be applied at temperatures sensibly higher than those which are compatible 219 with aliphatic amine-cured resins [12, p.168]. The completion of the curing pro-220 cess and the high thermal stability that is typical of amine-cured epoxy resins 221 are likely responsible for the increase in mechanical properties (elastic modulus and strength) that is observed in the epoxy-coated G-TRM composite materials that were treated at temperatures not exceeding 150 °C. At higher temper-224 ature, degradative phenomena are likely to outweigh the benefit conveyed by 225 post-curing. 226

#### 227 4.3. Failure analysis

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Fig.11 illustrates progression to the two typical failure modes: either fabric rapture (a) or fabric slippage inside the matrix (b). Although, generally, both of them occur in mixed proportion in all test groups, fabric failure is far more

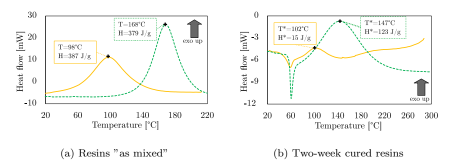


Figure 10: Differential scanning calorimetry (DSC) of EW resin (solid, yellow) and ER resin (dashed, green) right after mixing (a) and after two-week curing (b)

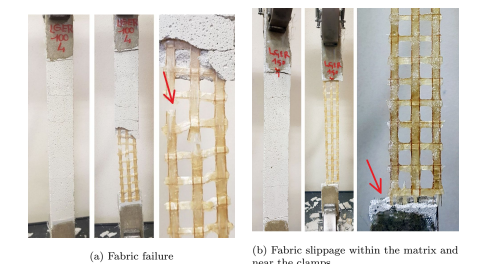
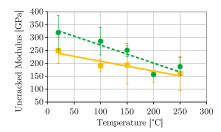


Figure 11: Typical failure modes observed for all groups

near the clamps



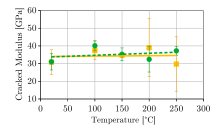


Figure 12: Uncracked (left) and cracked (right) modulus as a function of the exposure temperature for ER (circles, green) and EW (squares, yellow).  $\pm 1$  standard deviation bands and linear curve-fits are also given

frequent in the control group, while fabric slippage prevails in the specimens exposed to high temperature.

#### 5. Discussion

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Fig.12 illustrates the effect of temperature on the uncracked modulus  $E_f^*$ and on the cracked modulus  $E_f$  in the EW and in the ER group (see [14, 1] for the details of moduli definition and evaluation). It may be observed that temperature exposure strongly impairs the uncracked modulus  $E_f^*$ , while the cracked modulus  $E_f$  remains statistically unaltered. This is compatible with the expectation that temperature affects the coating performance, whose bearing is mostly relevant when the matrix is still collaborating with the fabric and thereby uncracked. Furthermore, although ER coating performs significantly better than EW, line fitting suggests that this advantage decreases with temperature until equal performance is met at T = 250 °C. Conversely, the cracked modulus  $E_f$  reflects the modulus of the glass fabric, which is little affected by temperature. In fact, the cracked modulus is about the same across all groups. Transition points (TPs) conventionally mark a sudden stiffness loss and a regime shift, in light of the fact that the cracked matrix ceases to contribute to the composite rigidity. Their location is shown in Fig.13 at different temperatures. This figure indicates that the transition stress nearly halves on high temperature exposure, irrespectively of the temperature value, in the EW group. Conversely, the ER group presents a similar transition point location for ER-100

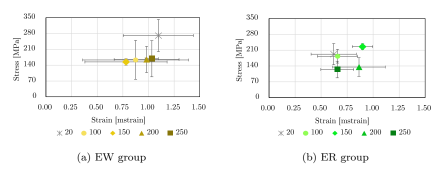


Figure 13: Mean transition point location and  $\pm 1$  standard deviation bars as a function of temperature for all test groups

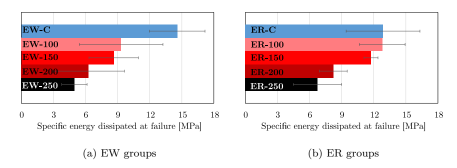


Figure 14: Mean specific energy dissipated at failure W and  $\pm 1$  standard deviation bars for the control (C, blue) and the heat treated groups at 100 °C (light red), 150 °C (red), 200 °C (dark red) and 250 °C (black).

and, remarkably, higher transition stress and strain for ER-150, with respect to the control group. Temperature adverse influence starts to manifest itself at 200 °C, when stress is impaired (-37.5%) and yet strain is still higher than in the control group. Finally, at 250 °C, strain drops and it reaches the value for the control group. It is concluded that temperature generally decreases transition stress but it may improve transition strain.

Comparison in terms of specific (per unit fabric volume) dissipated energy W is carried out in Fig.14. It appears that high temperature impairs energy dissipation in all test groups, with the possible exception of ER-100 which behaves similarly to the relevant control group. Remarkably, both coatings decay with an almost identical power-law rule, as illustrated in Fig.15 in terms of normalized quantities with respect to the ambient conditions. This finding is

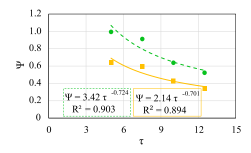


Figure 15: Normalized specific energy dissipated at failure  $\Psi=W/W_0$  against normalized exposure temperature  $\tau=T/T_0$  for ER (circles, green) and EW (squares, yellow) and power-law curve fit.  $T_0=20\,^{\circ}\mathrm{C}$  is the ambient temperature and  $W_0=W(T_0)$  is the corresponding dissipated energy.

compatible with a cumulative Weibull distribution for the relaxing and breaking
of the intermolecular bonds in the resin, as described in the model proposed by
Mahieux et al. [17], and it suggests that mechanical performance is indeed impaired by the mechanism of resin degradation. Conversely, hyperbolic tangent
models, as in [11, Eq.(5)], do not seem to fit well experimental data.

Fig.16 presents the behaviour of the relative ductility across all test groups 269 against temperature at different fraction of the UTS. Relative ductility is expressed as the ratio of the mean group strain  $\epsilon_i(f)$ ,  $i \in \{ER,EW\}$  over the mean 271 control group strain  $\epsilon_C(f)$ , when specimens are subjected to a traction force f 272 which is a fraction of  $f_{u_i}$ ,  $i \in \{ER,EW\}$ , that is the UTS for the relevant group. 273 It is clearly seen that the higher the temperature of conditioning, the more brittle specimens behave, with the single exception of 20% loading, see Fig.16(a). 275 However, for any temperature and loading fraction, ER outperforms EW in a 276 statistically significant manner. It is worth emphasizing that the EW group is 277 connected to superior energy dissipation capability in the control group, as com-278 pared to the ER group. This advantage at ambient temperature can be traced back to the EW coating layer being significantly thinner [19]. Therefore, it may 280 be argued that coating thickness is unfavourable in terms of ambient temper-281 ature mechanical performance, yet it is advantageous when high temperature 282 exposure is envisaged. 283

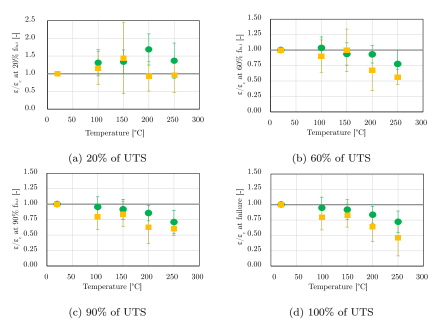


Figure 16: Ratio of the group mean elongation  $\epsilon$  to the relevant control group elongation  $\epsilon_C$  at a fraction of the corresponding UTS as a function of temperature for ER (circles, green) and EW (squares, yellow)

# 284 6. Conclusions

This works reports on the influence of the epoxy coating thermo-physical properties on the mechanical performance of AR-glass textile reinforced mortar (TRM) after exposure to high temperature. As in [19], two epoxy coatings are considered, which differ by the hardening agent alone. Nonetheless, this difference brings about distinct thermo-physical properties. Since focus is set on the epoxy coating, exposure temperatures are limited to 250 °C to prevent thermal effects from extending to the lime mortar and, eventually, to the glass fabric. Mechanical performance is assessed according to AC434 through uniaxial tensile tests of rectangular coupons. The effect of temperature exposure in terms of first cracking strength and strain, ultimate strength and elongation, cracked and uncracked moduli, transition point location and energy dissipation capability is illustrated. It is found that temperature exposure may increase strength at the expense of ductility, and this outcome parallels similar findings

obtained in the context of pull-out [8] and tensile [25] tests. DSC analysis reveals that temperature exposure may trigger competing processes: on the 299 one side further cross-linking is favoured in a post-curing process, on the other side thermal degradation occurs. The final outcome strongly depends on the 301 considered epoxy coating and its post-curing capability. Indeed, in contrast to 302 the findings reported in [9], mild degradation is documented, especially when 303 compared to FRP systems. Temperature induces a monotonic decay in the 304 energy dissipation capability and, remarkably, the decay law, that is the same for both coatings, complies with a cumulative Weibull distribution (power-law rule). 306 This behaviour is typical of models accounting for the relaxing and breaking of 307 molecular bonds in resins, as in [17, 11]. Therefore, this observation supports 308 the understanding that the resin degradation mechanism at the fabric-to-matrix interface governs mechanical performance for both coatings.

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# 317 Declaration of interest

Declarations of interest: none.

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