

Low energy pre-blended mortars: Part 2 – Production and characterisation of mortars using a novel lime drying technique

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1	Low Energy Pre-blended Mortars: Part 2 – Production and characterisation of
2	mortars using a novel lime drying technique
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11	
12	Abstract
13	
14	The presence of free water in mortars destined for silo or bagged storage can lead to the
15	degradation of the binder phase. Such water may be present as a result of using wet,
16	as-delivered sand or as a consequence of prior processes such as de-activation of
17	Roman cement. Thus, water must be removed from the system prior to storage. Part 1
18	of this paper describes the control of a technique by which quicklime is added to the wet
19	system which principally dries it by both slaking the quicklime and evaporation as a
20	consequence of the exothermic slaking reaction. Two examples of mortars are
21	presented in which excess water is removed from the system by the inclusion of
22	quicklime. In the first, the water is present in the as-delivered sand and the binder is a
23	combination of the slaked lime and ggbs. In the second, the water remains after pre-
24	hydration of a Roman cement which is a process to retard its rapid setting

25	characteristics. It is shown that optimally dried mortars are not subject to degradation
26	following storage of both mortar types.

28 **Keywords:** mortar, sand drying, ggbs, slaked lime, storage, Roman cement

29

30 **1** Introduction

31

32 The first part of this paper [1] described a technique by which the addition of quicklime to 33 wet sand could be controlled to produce Formulated Lime mortars which are dry and 34 suitable for long-term storage in silos or bags prior to final wet mortar production. The 35 principal process-control factors to achieve optimal drying have been identified as lime 36 addition based upon a fraction of the stoichiometric requirement for complete slaking of 37 the quicklime (i.e. 3.113 g of CaO being required to combine with 1 g water), free 38 moisture content of the sand and mixing time of the combined sand and guicklime blend 39 followed by a 24 hour "rest period". The philosophy can be equally applied to other 40 factory produced mortars in which free water may remain following the formulation 41 process.

42

A fundamental feature of this concept is that water which is not contained within the pore structure of the sand, i.e. free water, rather than the total water content is the one which must be controlled. In this scenario the datum state of the aggregate is its saturated surface dry (SSD) condition in which the sand pores are completely filled whilst the free moisture is zero. Trials of mortars produced with natural hydraulic lime (NHL 3.5) showed that mortars in which the binder was mixed with SSD sand and stored for 8

weeks in a sealed box exhibited no degradation when compared to companion mortars
which were immediately produced without the storage period.

51

52 This part of the paper describes the application of the technique in the production of two 53 types of mortar. The first are Formulated Lime mortars, M1 and M5, produced with the 54 addition of ggbs as the hydraulic phase. Formulated Limes are specified in BS EN 459-55 1:2010 [2] as being limes with hydraulic properties and comprising lime and added 56 material with hydraulic and/or pozzolanic properties whilst the strength classes M1 and 57 M5 (compressive strengths of 1 MPa and 5 MPa respectively) are specified in BS EN 58 998-2:2010 [3]. The mix design process has to ensure that the guicklime addition is 59 sufficient to yield both the dry mortar for storage and also the correct amount of ggbs, 60 principally the ggbs/slaked lime (CH) ratio, to yield the desired mortar type whilst 61 maintaining the required volumetric proportions of binder to aggregate. The properties of 62 mortars made immediately after addition of the hydraulic component are compared to 63 those in which the dry blended mortar was stored for 10 weeks prior to mortar 64 production. The second application is in the production of De-Activated Roman Cement 65 (DARC) mortars suitable for long-term storage. A characteristic of Roman cements is 66 their rapid setting such that retardation is required for the production of practical mortars. 67 A workable life of 1 - 2 hours has been specified for render mortars [4] for which a pre-68 hydration process has been developed [5]. In essence, this process involves adding an 69 initial controlled amount of water to the Roman cement such that sufficient reaction takes 70 place to retard the setting of the mortar when the fresh mortar is subsequently 71 manufactured. During that study it was observed that DARC mortars stored for 1 - 1272 weeks exhibited some degradation during storage, thought to be due to the presence of 73 residual water following the de-activation process. Thus, the situation is akin to that of 74 the presence of excess water in the Formulated Lime mortars which needs to be

75	removed, although in this case the water results from a factory-based process rather
76	than from as-delivered materials. Mortars with the addition of quicklime following the de-
77	activation stage are compared with samples which were oven-dried at this point as well
78	as control samples.
79	
80	Part A: Formulated Lime Mortars
81	
82	2 Materials and Methods
83	
84	2.1 Materials
85	
86	A siliceous sharp sand was used throughout and was oven dried to constant weight at
87	110°C prior to use. Two different quicklimes, C1 and E, were used for the preparation of
88	the lime-dried mortars. The chemical and physical properties of these materials, together
89	with properties of the slaked limes, were reported in Part I of this study [1]; Table 1
90	summarises the key properties of the slaked limes and a commercial CL90 which has
91	been used as a control. ggbs was supplied by Civil & Marine Slag Cement from the
92	Frodingham plant, UK.
~ ^	

Lime	C1	Е	CL90
Free lime (% CaO)	70.9	68.7	71.3
Insolubles (wt%)	5.8	4.8	3.8
CaO (wt%)	0.3	0	0.9
Ca(OH) ₂ (wt%)	93.2	90.8	93.0
CaCO ₃ (wt%)	2.9	6.9	2.3
A _{BET} (m²/g)	15.19	18.16	11.26
V _{TOT} (cm ³ /g)	0.098	0.109	0.072
Bulk Density (kg/m ³)	585	559	575

Table 1: Properties of limes

2.2 Mortar Production

2.2.1 Mix design of lime-dried mortars





Fig 1: Example of mix design for optimum mortars using lime E.

102 Figure 5 of ref [1] shows that the optimum stoichiometric ratio for quicklime addition to dry the sand decreases with increases in either the sand moisture content or the mixing 103 104 time of the wet sand and quicklime. This is represented for lime E in Figure 1 by the 3 105 linear relationships. The design of the lime-dried mortar formulations was intended to 106 produce mortars of M1 and M5 classification according to BS EN 998-2 [3]. The 107 approximate ggbs/CH ratio (by weight) required to achieve the stated classification was 108 0.2 and 1.0 respectively where the mortars were produced to a common flow of 170 mm 109 [6]. Obviously, the precise ratio required to yield a specified strength for any mortar will 110 also be related to the purity of the quicklime, grading of the sand as it affects workability, 111 the sand density and the density of the $Ca(OH)_2$ (also referred to by its cement chemistry 112 notation of CH) generated during the drying process as they affect volumetric mix 113 proportions. Once a sand:binder ratio has been selected it is possible to calculate how 114 much Ca(OH)₂ is required for a specified ggbs/CH ratio. Hence, knowing how much 115 quicklime would be required to chemically combine with all of the free water for a given 116 free moisture content of the sand, the stoichiometric ratio of guicklime addition may be 117 calculated which generates the correct amount of Ca(OH)₂ for the specified ggbs/CH 118 ratio, after accounting for the purity of the quicklime. For instance, for a particular mortar 119 batch size and mix proportions, if 1381 g of Ca(OH)₂ is required to yield the desired 120 ggbs/CH (equivalent to 1045 g of CaO), whilst theoretically requiring 1771 g of CaO to 121 chemically combine with the free water in the sand, this would represent a stoichiometric 122 ratio of 59%. In this procedure the density of the ggbs and the CH have been measured 123 but no allowance was made of any interactions between the two phases which might 124 affect their packing efficiency.

125

126 The associated values of stoichiometric ratio and sand free moisture content for both 127 ggbs/CH ratios have been superimposed on Figure 1 and the intersection of the

relationships indicates the optimum drying conditions for each mortar; the 4 mortars using quicklime E detailed in Table 2 are indicated on Figure 1. The same approach was adopted for mortars using quicklime C1 In accordance with the terminology previously described [1] all mortars are considered optimally dried. All mortars were produced at a constant sand:binder ratio of 2.25:1 by volume.

Code	Lime type	Stoich ratio	Sand m/c	Mix time	ggbs/CH ratio	w/b ratio
		(%)	(%)	(min)		
M1-1	Е	58.87	5.69	15	0.2	1.35
M1-2	Е	62.62	5.35	10	0.2	1.36
M1-3	Е	69.72	4.81	5	0.2	1.36
M1-4	C1	61.20	5.71	15	0.2	1.37
M1-5	CL90	~	~	~	0.2	1.41
M5-1	Е	60.9	4.03	15	1	1.03
M5-4	C1	61.2	4.08	15	1	1.05
M5-5	CL90		~	~	1	1.06

¹³³

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Table 2: Mix design
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134

The experimental programme was designed to determine the effect of various parameters of the lime-drying process on the physical properties of the resulting hardened mortars. The formulations allow assessment of the influence of the drying process, the drying time and the quicklime source; mortars using CL90 provide control data.

140

141 **2.2.2 Preparation of lime-dried mortars**

143 Ten kg batches of sand were prepared containing known amounts of free water (wt.% 144 basis). The sand was then mixed in a Hobart mixer with appropriate quantities of 145 quicklime for the specified period of time (i.e. 5, 10 or 15 min). Following mixing, the 146 lime-dried materials were stored in sealed containers for 24 hours to ensure complete hydration of the quicklime; this step was included after previous observations indicated 147 148 the presence of free water in the blend at the completion of the initial mixing [1]. The 149 required quantity of ggbs was then added in order to provide the desired mix 150 proportions. At this stage, the dry mortar mix was divided into two equal portions. One 151 portion was stored in a sealed plastic container and retained under laboratory conditions 152 (20°C, 50% RH) for 10 weeks; the other portion of the mix was prepared immediately. 153 These mortars have been termed "stored" and "fresh" respectively.

154

155 **2.2.3 Production of plastic mortars**

156

Lime dried mortars were produced by adding mix water to the "dry mortar" whereas the control mortars were produced by adding the water to a blend of CL90 and SSD sand. All plastic mortars were produced in a Hobart mixer with a mixing time of 15 minutes. Appropriate quantities of water were added to give a flow table value [7] of 170mm (+/-5mm). This value was considered to provide materials with workability consistent with that observed in typical construction practice [8]. The required free water:binder ratio to achieve the specified flow value for each mortar is displayed in Table 2.

164

165 **2.2.4 Curing of mortars**

166

Following preparation, the plastic mortars were cast into steel moulds of the required geometries (see section 2.3). The samples were initially cured within the moulds for 3

169 days at 20°C and covered by a polythene sheet to minimise evaporation. The moulds 170 were then stripped and the samples cured for a further 4 days under the same 171 conditions. Unless stated otherwise, extended curing was then carried out at 20°C (\pm 172 2°C) and 65% RH (\pm 5%), hereafter referred to as air curing. The level of CO₂ in the 173 curing room was assumed to be the standard atmospheric concentration (0.033 \pm 174 0.001% by volume). For comparison, some samples (for strength testing only) were 175 subject to extended curing under water, also at 20°C.

176

177 **2.3 Analytical Methodology**

178

179 **2.3.1 Compressive and Flexural Strength**

180

181 The test specimens were prepared in steel moulds of 40 x 40 x 160 mm dimensions. 182 Mortar was placed in 2 layers and compacted using a vibration table. The three-point 183 flexural and compressive strengths of the hardened mortars were determined using an 184 Instron 4206. Crosshead speeds of 0.5 mm/min and 1 mm/min were used for flexural 185 and compressive testing respectively. Measurement of compressive strength was 186 carried out at 7, 28, 91 and 365 days. For air-cured samples, flexural strengths were 187 determined at 7 and 91 days whereas water-cured materials were tested at 28 and 91 188 davs.

189

2.3.2 Density and Porosity

191

The bulk density and total open porosity of the hardened mortars were assessed in accordance with BS EN 1015-10 [9] and BS EN 1936 [10] respectively (vacuum saturation method). The reported values are the mean of three individual samples.

195 **2.3.3 Sorptivity**

196

The water sorptivity of the hardened mortars was determined according to the methodology described in detail by Hall [11]. The test specimens were prepared in 70 mm steel cubes and cured as described in section 2.2.4. Prior to testing, the samples were conditioned at 60°C to constant weight in an atmosphere circulated over silica gel and soda lime. The sorptivity was measured through the moulded bottom surface; the vertical faces were sealed by the application of water-resistant tape. The reported values represent the mean of three individual samples.

204

205 **2.3.4 Water Vapour Permeability**

206

207 The water vapour permeability of the hardened mortars was determined using a 208 modified version of the methodology described in BS EN 1015-19 [12]. The test 209 specimens were cast in circular steel moulds, producing specimens of 180 mm diameter 210 and 20 mm thickness. The mortar discs were then subjected to the curing regime 211 specified in section 2.2.4. After curing, the samples were prepared for analysis at 60°C in 212 an atmosphere circulated over silica gel and soda lime. The samples were then placed 213 in stainless steel test cups containing a saturated solution of potassium nitrate and 214 sealed in-situ with molten paraffin wax. The air gap between the base of the samples 215 and the solution was 12 mm (± 2 mm). The KNO₃ solution generates a relative humidity 216 within the air gap of 93.2% at 20°C. The samples were then placed in a fan-assisted 217 storage chamber maintained at 20°C (±2°C) and 50% RH (±5%) and weighed every 24 218 hours until steady state vapour transmission was achieved. The reported values 219 represent the mean of three individual samples.

220

221 **2.3.5 Mercury Intrusion Porosimetry (MIP)**

222

223 The pore size distributions of the mortars were determined using MIP at an age of 91 224 days (air cured). The measurements were carried out with a Micromeritics 9400 series instrument over a range of pressures between 3.9x10⁻³ and 410 MPa. Assuming a 225 contact angle of 140° and a mercury surface tension of 484x10⁻³ N/m, pore diameters 226 ranging from 375 μ m to 3.8x10⁻³ μ m were characterized. Specimens with an approximate 227 volume of 2 cm³ were cut from the mortar prisms using a low speed saw. Prior to 228 229 analysis, the samples were dried to constant weight at 60°C in an atmosphere circulated 230 over silica gel and soda lime.

231

3. Properties of "fresh" mortars

233

3.1 Pore structure

235

Values of the total open porosity and dry bulk density are shown in Table 3. The general trend observed for all mortars is a slight decrease in porosity, allied with an increase in bulk density, as the curing time was extended, and is more marked in the M1 mortars. The M5 series of mortars show higher density and lower porosity than the M1 series, as would be expected from the higher ggbs/CH ratio. The difference is more marked in mortars produced using either quicklime C1 or the CL90.

Mortar	Open Porosity (%)			Dry Bu	Ik Densit	y (kg/m³)
	28 d	91 d	365 d	28 d	91 d	365 d
M1_1	32.2	31.6	31.0	1757	1781	1791
IVI I - I	32.7	31.8	31.4	1689	1717	1729
M1-2	31.8	31.4	31.0	1769	1794	1799
WI 1-2	32.9	31.7	31.5	1684	1719	1735
M1-3	32.1	31.4	30.8	1761	1795	1797
WI 1-5	32.8	31.9	31.4	1689	1714	1737
M1-4	33.5	32.8	32.3	1720	1749	1767
1011-4	34.2	33.1	32.6	1660	1700	1709
M1-5	34.9	34.2	33.7	1693	1686	1701
WI 1-5	35.1	34.2	33.6	1651	1679	1700
M5-1	29.3	29.3	29.1	1767	1805	1805
10-1	29.9	28.5	27.8	1733	1746	1779
M5-4	29.4	29.4	29.1	1755	1777	1784
1413-4	30.1	29.1	27.7	1729	1741	1775
M5-5	29.5	29.8	29.3	1745	1759	1767
1410-0	30.6	29.0	28.2	1721	1733	1766

242Table 3: Porosity and density of mortars at ages of 28, 91 and 365 days (air curing in243Normal font, water curing in Italics).

Within the M1 range of mortars M1-1, M1-2 and M1-3 possess the same porosity at a given age and curing condition; all these mortars were dried with the same quicklime E, the only difference being in the process parameters of stoichiometric ratio and mixing time. The influence of lime type may be seen in the higher porosities of M1-4 (quicklime C1) and M1-5 (CL90). However, any influence of lime type in the M5 mortars (M5-1, M54 and M5-5) is reduced to a minimum, probably as a function of the reduced lime content
of these mortars.

252

The influence of curing regime is small for the M1 mortars although water curing yields the slightly higher porosity at all ages. In contrast, water-cured M5 show higher porosity at 28 days but lower porosity at 365 days, presumably reflecting additional hydraulic activity at later ages.

257

258 The pore size distributions of the mortars (91 days of air curing) are shown in Figure 2. 259 The data highlight significant differences in the pore size distributions between the M1 260 (Fig 2a) and M5 (Fig 2b) mortar series. The lime-rich M1 series contain a large proportion of pores in 2 distinct zones, i.e. $10 - 18 \mu m$ and $0.2 - 0.6 \mu m$. Mortars M1-4 261 262 and M1-5 exhibit similar pore structures; both mortars are based upon lime from the 263 same source. Mortar M1-1 shows a slightly lower threshold pore diameter and a lower 264 volume of the coarsest pores which accompanies the lowest open porosity (Table 3). In 265 contrast, for the M5 mortars, the bulk of the porosity is made up of pores in the region of 266 0.05 µm diameter. This is a function of the higher ggbs/CH ratio and lower w/b ratio of 267 the M5 series. As with the M1 mortars, the pore distributions of M5 mortars using 268 quicklime C1 and CL90 are similar.



269 Fig 2: Pore size distribution of (a) M1 and (b) M5 mortars after 91 days of air-curing.

A similar pattern of small differences in open porosity accompanied by marked
differences in pore size distribution has been previously reported by Arandigoyen and
Alvarez [13] for cement-lime mortars.

274

3.2 Compressive and flexural strength

276

277 **3.2.1 M1 mortar formulations**

278

279 The data in Figures 3a and 3b show the evolution of flexural and compressive strength 280 of the M1 mortars subjected to air curing. Under flexural load, mixes M1-1 to M1-5 yield 281 similar strengths at 7 days. All mortars show a considerable and similar gain in flexural 282 strength up to 91 days (all significance testing used the student t-test conducted at 95% 283 confidence). Under compression, mortars M1-1 to M1-5 display essentially similar initial 284 strength development up to 28 days. Thereafter, mortars M1-1, M1-2 and M1-3 show a 285 further strength increase up to 91 days with no further change between 91 and 365 286 days. In contrast, mortars M1-4 and M1-5 show no gain in compressive strength after 28 287 days and, consequently, these materials are significantly weaker at the later ages.



Fig 3: Strengths of M1 mortars – (a) flexural strength air curing, (b) compressive strength air curing, (c) flexural strength water curing, (d) compressive strength water curing. Error bars represent ±1 Standard Deviation.

292 Figures 3c and 3d display the influence of water curing on the strength development of 293 the mortars M1-1 to M1-5. In flexure, mortars M1-1, M1-2 and M1-3 exhibit similar 294 strengths at 91 days with mortars M1-4 and M1-5 also being similar but weaker than the 295 other mortars. With the exception of M1-4, the strength of all mortars is similar at 91 296 days for both curing conditions. Up to an age of 28 days, all mortars show similar 297 compressive strengths and there is little difference in the compressive strengths of 298 samples subjected to the different curing conditions. However, at ages of 91 and 365 299 days water curing generally generates higher strength (typically 20 - 40%) reflecting the 300 extra potential for hydraulic reactions in the lime-ggbs system. The exception is mortar 301 M1-3 which exhibited a high variability of strength under air curing at these ages, so influencing the statistical comparison; it is not believed that the apparent differenceexhibited by this mortar formulation is genuine.

304

305 **3.2.2 M5 mortar formulations**

306

The data in Figures 4a and 4b show the evolution of flexural and compressive strength of the M5 mortars. There is little influence of lime type in the M5 mortars. As expected the highest strengths are registered by water cured samples such that by an age of 1 year a strength enhancement in the range 130 – 140% is observed. Whilst the latter curing yields continuous strength increase up to an age of 1 year, air cured samples indicate a maximum strength at either 28 or 91 days with a subsequent reduction at an age of 1 year; this was not observed in the M1 mortars.



Fig 4: Strengths of M5 mortars – (a) flexural strength, (b) compressive strength. Error
 bars represent ±1 Standard Deviation.

316

The influence of curing regime on long term development of alkali-activated slag concrete (sodium silicate and calcium hydroxide activation) [14] and of calcium hydroxide activated slag mortars [15] has been previously observed. The strength reduction was explained by the high shrinkage and micro-cracking associated with this binder system. It may be observed that the absence of strength reduction is seen in the 322 lime-rich M1 mortars and, whilst any correlation is purely speculative, the strength 323 reduction in the M5 and calcium hydroxide activated slag mortars [15] occurs in mortars 324 with a larger hydraulic component and less lime.

325

Both compressive and flexural strengths have been measured for all mortars and curing conditions at 2 ages. The ratio of compressive:flexural strength as a function of compressive strength is shown in Figure 5. As expected the ratio increases as the compressive strength increases reflecting a more brittle material. There is a certain degree of scatter, particularly at lower strengths; a correlation coefficient of 0.872 is achieved.



332

Fig 5: Ratio of compressive:flexural strength as a function of compressive strength forM1 and M5 mortars.

335

Both M1 and M5 mortars generally achieve their strength designation at an age between 28 and 91 days of water curing although it is likely the time is towards the lower end of this range for the M5 mortars.

339 **3.3 Moisture transport**

340

341 The specification of mortars is dependent upon their end-use. In the case of restoration 342 works on historic buildings their breathability is an important property. However, this is a 343 topic with differing philosophical approaches adopted by different restorers. One 344 approach, as codified in BS EN 998-1:2010 [16], may be summarized as one in which 345 the movement of water is to be minimised where-as the movement of water vapour is to 346 be encouraged. This approach frequently requires the use of hydrophobic treatments 347 within the mortar system. Another viewpoint is that this approach only transfers the 348 original problems to other areas of the structure which have not been treated with such a 349 mortar [e.g. 17, 18]. It is argued that since water transport is much quicker than vapour 350 transport both parameters are important.

351

In the current work, breathability was assessed by sorptivity and water vapour
permeability (WVP) measurements following air curing and the results are presented in
Figures 6 and 7 respectively.



355





358

Fig 7: WVP of M1 and M5 mortars.

359

360 It is apparent that the sorptivity and WVP of the M5 mortars is less than that of the M1 361 mortars reflecting their finer pore structure. Although the average values suggest that all 362 mortars show a variation in sorptivity with age, the variability within each data set is such 363 that there is no statistical difference between mortars. By way of contrast, the WVP of 364 the M1 mortars generally increases with age although that of the M1-5 mortar only 365 marginally fails the significance test. There is no influence of lime source on the 366 sorptivity or WVP of the M1 mortars at ages either 91 or 365 days unlike that observed 367 for compressive strength, nor on the sorptivity or WVP of the M5 mortars at 91 days 368 which is the only test age with data for all of this group of mortars.

369

Both M1 and M5 mortars exceed a commonly accepted value of 4 Kg/m²/hr^{0.5} for the sorptivity, or Water Absorption Coefficient, for restoration mortars for renders [4]. Obviously, neither mortar type complies with the low water absorption criterion of BS EN 998-1 although the M1 mortars do meet the requirement for WVP (μ >15).

The lower "permeability" of the M5 mortars is reflected in their reduced rate of carbonation as measured by depths of complete carbonation using the phenolphthalein test (Table 4).

Mortar	28d	91d	365d
M1	2 - 3	9 - 10	Complete
M5	1 - 3	6 - 7	13 - 15

378

Table 4: Carbonation depths (mm) of M1 and M5 mortars

379

380 **3.4 Discussion**

381

An influence of lime type has been observed in strength performance but not in moisture transport. It is possible that this may be a function of the greater variability within the latter determinations and had a larger number of samples been tested at each stage a significant difference might have been obtained. The following discussion will focus on the strength data.

387

388 The similar performance of the M1-1, M1-2 and M1-3 mortars is to be expected as the 389 materials are of essentially identical composition with the key parameters of lime source 390 and ggbs/CH & w/b ratios being common. The data provides confirmation that 391 manipulation of the lime drying process (quicklime stoichiometric ratio and mixing time) 392 allows the production of identical mortars from quarried aggregates containing different 393 amounts of free water. Likewise, the performance of mortars M1-4 and M1-5 may be 394 similarly explained; despite M1-4 being a lime-dried mortar and M1-5 comprising a 395 commercial CL90, the source of quicklime in both cases is the same.

374

397 It was previously shown that lime E yielded higher long term strengths than lime C1 in 398 the M1 mortars but not in the M5 mortars (sections 3.2.1 and 3.2.2). The differences in 399 BET surface area (Table 1) do not account for the different performances; however, the 400 greater strength of the mortars produced with lime E may reflect the higher CaCO₃ 401 content of the slaked lime. In this case, the lime-rich M1 mortars contain a larger quantity 402 of CaCO₃ than do the M5 mortars. Studies of the use of CaCO₃ as an addition to 403 Portland cement [19] show that fine $CaCO_3$ can be both a reactive component as well as 404 a filler. Additional hydration within the AF_m phases is observed at low carbonate 405 additions and is accompanied by reduced porosity. A study of the inclusion of metakaolin 406 and limestone in Portland cement mortars has shown that the carbonate reacts with the 407 metakaolin to produce AF_m phases [20]. The identification of AF_m in alkali-activated 408 cements is uncertain and may be depend upon detection technique and the particular 409 alkali utilized [e.g. 21, 22]. Further, whilst the addition of 1-3% fine calcite has been 410 found to yield a small increase in the early age strength of an alkali-activated cement, no 411 or even a negative impact occurred at later ages [19]. Little modern research has been 412 undertaken on the lime-ggbs system and certainly not in the low slag contents currently 413 being considered and further fundamental research is required to examine the role of 414 $CaCO_3$ within this system.

415

In contrast to the M1 mortars, the strength development of M5-1, M5-4 and M5-5 mortars appears unaffected by the different limes used in their manufacture. The reasons for this difference are unclear but the discrepancy indicates that the properties of the lime are of greater significance in mortars with higher lime content. It seems reasonable to assume that at low ggbs/lime ratios, the development of the mortar microstructure would be more significantly influenced by changes in lime properties than

in mortars where the lime content is much reduced. In cement-lime-sand mortar systems, Sebaibi *et al.* [24] stated that 'it is necessary to have a high lime substitution percentage to influence the microstructure of the mortar'; however, the PC/lime ratios used in their mortar formulations were far higher than the ggbs/lime ratios employed in the current work.

427

- 428 **4** The effect of storage on lime-dried mortar mixes
- 429
- 430 **4.1 Optimally dried mortars**
- 431

432 The data shows that the performance of lime-dried mortars is very similar to those 433 produced using commercial CL90 hydrated lime. However, one of the essential 434 requirements of the lime-drying process is the ability to produce mortars that can be 435 stored in silos or bags without significant degradation prior to use. The lime-drying 436 process is unlikely to remove water absorbed within the pores of the aggregate unless 437 excessive heat is generated during the slaking process. As a pre-cursor to work on lime-438 dried mortars a range of mortars was produced using NHL 3.5 as the binder phase in 439 which the sand was pre-prepared at various moisture states between oven dry and 440 saturated surface dry. They were blended in the volume ratio 2.25:1 and stored for 8 441 weeks prior to mixing into mortars of common flow. There was no statistical difference 442 (95% confidence limits) between mortar strengths at ages up to 2 years indicating no 443 influence of absorbed water on mortar performance. Hence, a series of lime-dried 444 mortars was produced following a 10-week storage (see section 2.2.2) in which the 445 expectation was that any influence of storage would be a function of free water rather 446 than absorbed water.



447 Fig 8: Relative strengths of stored and fresh mortars; (a) air cured, (b) water cured.

449 Figure 8 shows the relative compressive strength of the M1 and M5 series of "stored" 450 mortars (both water and air-cured) in comparison to the "freshly" prepared materials 451 described in section 3.2. It is apparent that there is a difference in performance between 452 the M1 and M5 mortars. The M5 "stored" mortars are generally similar to their "fresh" 453 counterparts at all ages except 365 days where the air-cured "stored" mortars lose 454 strength at a more rapid rate than the "fresh" mortars (see Fig 8a). At 365 days the 455 "stored" mortars are some 77% of the 91 day strengths in contrast to 87% for the "fresh" 456 mortars. In contrast, the strengths of the M1 "stored" mortars are considerably greater 457 than those of the fresh materials at ages beyond 7 days under both air and water curing. 458 The strength increases are accompanied by an increase in bulk density and a decrease 459 in the porosity in the stored mixes (Table 5).

Mortar	Open porosity (%)		Density	′ (kg/m³)
	Fresh	Stored	Fresh	Stored
M1-1	31.6	30.6	1781	1809
M1-2	31.4	30.6	1794	1808
M1-3	31.4	30.2	1795	1818
M1-4	32.8	30.9	1749	1803
M1-5	34.2	32.6	1686	1766

Table 5: Influence of storage on M1 mortars

461

401

462 Only a very limited study was undertaken on M1 mortars (M1-1, M1-4 & M1-5) to 463 examine the influence of storage upon the moisture transport properties at an age of 91 464 days curing in air. Whilst strength was affected by storage there is no significant 465 influence on moisture transport.

466

467 The precise reason for the observed differences between the M5 and M1 mortar series 468 is unclear but appears to be related to changes in the water demand of the plastic 469 mortars following storage. Table 6 shows the w/b ratios required to achieve the specified 470 flow values during mortar manufacture. For the M1 series, the data show a marked 471 reduction in water demand after storage whereas comparatively little change is observed 472 for the M5 series. It seems reasonable to assume that the reduction in water demand is 473 a result of changes in the properties of the calcium hydroxide during storage. These 474 changes were discussed in detail elsewhere [1] and include decreases in surface area 475 and possibly a small increase in the quantity of calcium carbonate within the lime; see 476 Tables 8 and 9 of ref [1]. For a given lime, it seems likely that a reduction in surface area 477 would lead to a lower water demand in the plastic mortars and, as discussed previously, 478 the presence of calcium carbonate leads to enhanced strength in lime mortars. 479 Furthermore, the high proportion of lime in the M1 mixes suggests that these materials 480 would be more sensitive to changes in lime composition than the slag-rich M5 series. 481 Despite the increasing use of pre-mixed lime mortars in UK construction practice, there 482 is no evidence within the literature of research to determine potential changes in the 483 properties of these materials following extended periods of silo storage. In the current 484 paper, changes were not confined to mortars prepared with the lime-drying technique 485 (see mortar M1-5) and it seems reasonable to assume that other lime rich mortars may 486 also be subject to the same phenomenon. Hence, further research is recommended in 487 this area.

Mortar	w/b ratio		Change
	"Fresh"	"Stored"	(%)
M1-1	1.35	1.30	3.7
M1-2	1.36	1.31	3.7
M1-3	1.36	1.29	5.1
M1-4	1.37	1.31	4.4
M1-5	1.41	1.35	4.3
M5-1	1.03	1.01	1.9
M5-2	1.01	1.00	1.0
M5-3	0.99	0.98	1.0

488 Table 6: Influence of storage on w/b ratio to achieve common flow.

489

490 **4.2 Non-optimally dried mortars**

491

Three regions of drying have been previously defined [1]. Optimal drying is considered to

493 occur when the measured evaporation during the drying process equals the balance of

the original free moisture and that removed by chemical combination after accounting for the stoichiometric ratio of quicklime used and its purity. In contrast, sub-optimal drying does not remove sufficient water and super-optimal drying removes too much water leaving a residual amount of free lime - see ref [1] for a fuller discussion. The assumption was that sub-optimal drying would lead to degradation during storage whilst super-optimal drying would yield an unsound binder. The latter has previously been confirmed for storage periods of up to 4 weeks [1].

501

502 A series of 10 mortars was produced based upon the optimal processing criteria for the 503 M1-1 mortar. By reducing the stoichiometric ratio of quicklime sub-optimal mortars were 504 produced and by increasing it super-optimal mortars resulted. After lime-drying half the 505 mortar was used to produce "fresh" mortars whilst the remainder was stored in sealed 506 boxes for 10 weeks to produce "stored" mortars. All mortars were produced to a 507 common flow of 170 mm and subject to both air and water curing. Table 7 shows the key 508 data of stoichiometric ratio and the stoichiometric variation from optimal conditions for 509 each mortar together with the w/b ratio to achieve the desired flow. It can be seen that 510 the optimally dried mortar has the highest water demand of all mortars; the reason for 511 which is not immediately apparent.

Stoichiometric ratio	Stoichiometric variation	w	/b
(%)	(%)	"Fresh" "Stored"	
50.0	-9	1.18	1.17
54.9	-4	1.21	1.25
55.9	-3	1.21	1.26
56.9	-2	1.24	1.24
57.9	-1	1.28	1.23
58.9	Optimal	1.35	1.31
59.9	+1	1.26	1.22
60.9	+2	1.28	1.22
61.9	+3	1.28	1.20
67.0	+8	1.26	1.18

Table 7: Sub-optimal, optimal and super-optimal mortars

513

514 The water demand of the "stored" optimal and super-optimal mortars is less than their 515 "fresh" counterparts as previously observed in section 4.1. This might be explained by 516 the presence of unslaked lime in the "fresh" mortars placing a demand for water to slake 517 the quicklime during mortar production whilst this was air-slaked during the storage 518 period. In contrast, the trend for the sub-optimal mortars is less clear but, in general, 519 appears to be either unaffected or shows an increase in water demand which may be a 520 reflection of the generation of increased surface area by hydration of ggbs during 521 storage.

522

523 It is clear from the raw data that the use of super-optimal drying has not been 524 detrimental to strength development under either storage period or curing condition. This

525 would support the supposition that any free lime remaining after lime-drying has been converted to slaked lime during the storage period or the mortar mixing process. The 526 527 relative performance of each mortar, expressed as the ratio of the compressive strength 528 of the "stored" / "fresh" mortars at ages between 7 - 365 days is shown in Figure 9. 529 Under both curing conditions both the optimal and super-optimal mortars display a 530 strength enhancement following the 10 week storage period; the super-optimal +1% 531 mortar may be an anomaly. Whilst the sub-optimal -1% and -2% mortars subject to air 532 curing also show strength enhancement at all ages, the water cured samples register a 533 reduction in relative strength with age suggesting that the longer term hydration of the 534 ggbs may have been compromised during storage. The sub-optimal -3% mortar appears 535 to present a threshold in behaviour in that whilst storage has not affected the strength 536 neither is strength enhancement observed. As the stored mortars become wetter (-4% 537 and -9% variation) it is clear that they have degraded to the extent that the "stored" 538 mortars are substantially weaker at all ages, more so following water curing than air 539 curing.



540Fig 9: Influence of varying the quicklime addition on relative strength of "fresh" and541"stored" mortars subject to (a) air curing and (b) water curing.

549	5 Materials and Methods
548	
547	Part B – Roman cement mortars
546	
545	also minimizing the financial cost of using excessive quicklime.
544	a narrow range of quicklime addition in which the quality of mortar is maintained whilst
543	This data suggest that in the practical implementation of the lime-drying process there is

551 A Roman cement developed during the EU funded ROCARE project from marls sourced 552 in Gartenau, Austria, and a 0 - 4 mm carbonate sand were used. The cement was 553 manufactured by The Institute of Ceramics and Building Materials (MBM) in Krakow, 554 Poland; details may be found elsewhere [5]. Two mixes have been de-activated with 7% 555 de-activation water and 30 min storage prior to the formulation of the final mortar. The 556 deactivation water, expressed as a percentage of the cement weight, is first added to the 557 oven dry sand and mixed for 2 min at 62 rpm (a Hobart mixer was used for all mortar 558 production). Subsequently, the cement was added to the wet sand and the whole mixed 559 for a further 2 minutes at 62 rpm before being stored in an airtight box. At the end of the 560 30 minute de-activation storage time, the first mix was oven dried for 24 hours at 105°C 561 in an atmosphere circulated over silica gel and soda lime while the second mix was lime-562 dried by means of the addition of a pre-determined amount of quicklime. The moisture 563 content of the DARC mortar was determined as 0.63%. Based on previous experience 564 the lime content was specified as 76% stoichiometric ratio with a 5 minute mixing time. 565 Both mixes were then stored in airtight boxes for 4 weeks prior to formation into mortars. 566 A third mix (control mix) was prepared and stored for the same period of time with no 567 drying carried out.

At the end of the 4 week storage period, mortars were manufactured at a sand:cement ratio of 2.5:1 by volume and constant w/c ratio of 0.81. The mortars were produced by mixing the products of the previous drying processes in a Hobart mixer with water for 30 s at 62 rpm. At this time the mixer was stopped for 30 s during which the mortar adhering to the wall and bottom part of the bowl was removed by means of a rubber scraper and placed in the middle of the bowl. The mixing was then continued at 125 rpm for 8 minutes.

576

Flow and workable life were measured according to BS EN 1015-3:1999 [7] and BS EN
1015-9:1999 [25] respectively. Samples for strength determination were cast in 40 × 40
× 160 mm steel moulds in two layers and vibration compacted before being covered with
a polythene sheet. Mortar beams were de-moulded after 24 h and cured in water at 20°C
prior to testing at 28 days.

582

583 Differential Thermo-gravimetric (DTG) tests (a heating rate of 5°C per minute to 150°C 584 under a nitrogen atmosphere) were also performed on the oven dried mix and on the 585 control mix immediately after the end of the 30 minute storage period.

Mortar	ortar Flow Work Life		Comp Strength	Flex Strength
	(mm)	(min)	(MPa)	(MPa)
Control	170	>141	9.84	2.98
Oven Dried	155	~50	11.8	3.51
Lime Dried	140	~47	13.2	3.92

589

Table 8: Properties of Roman cement mortars

590

591 In Table 8 it is shown that both the oven dried and the lime-dried mortars exhibit a lower 592 fluidity than that of the control mortar, with the lowest flow being observed for the lime-593 dried mortar. This means that these mortars will require higher amounts of water to 594 achieve the desired workability. Whilst the higher water demand of the lime dried mortar 595 can be explained by the presence in this mix of calcium hydroxide formed in the reaction 596 of quicklime with de-activation water, it was not expected that the oven dried mix would 597 have a higher water demand than that of the control mix. In fact, in a previous study [5] it 598 was shown that during storage of a DARC mortar (i.e. the control mortar) the water 599 demand for the subsequent wet mortar was increased as the storage period was 600 prolonged. Whilst all 3 "dry" mortars in the current study were stored for 4 weeks, the 601 oven-dried mortar was conditioned to a lower free moisture content such it might have 602 been expected to perform similarly to a DARC mortar that had been subject to a 30 min 603 only storage period, i.e. a lower water demand or higher fluidity than that of the control 604 mortar. A possible explanation of this data is that the oven-drying causes the de-605 hydroxylation of the AF_m phases. In order to verify this hypothesis, DTG tests were 606 undertaken on the control mix (fresh) and on the oven-dried (OD) mix and results are shown in Figure 10. It is apparent that the use of oven drying has removed water from the system which may be accounted for by both the loss of evaporable water [26] and partial de-hydroxylation of the AF_m phases [27]. In this case, the reverse process (rehydroxylation) might take place during the early life of the constituted mortar and account for the increased demand for mixing water.



612

Fig 10: DTG curves of control (fresh) mix and oven dried (OD) mix (5°C per minute to

614

150°C).

Table 8 also shows that the workable life of the mortar produced with the control mix is at least 3 times longer than that produced with the oven dried and with the lime-dried mix. This can be in part attributed to the higher flow of the control mix mortar and in part to the longer storage under conditions of free moisture experienced by this mix, which has been shown to increase workable life [5]. Further study including the production and testing of mortars at constant flow is necessary to better understand the effect of the drying processes on workable life. Both compressive and flexural strength (28 days) are

higher for the dried mix mortars than for the control mix mortar, with the highest strengthobserved for the lime-dried mix mortar.

625

626 It is not possible to undertake a strict comparison between the degradation observed in 627 the stored DARC mortars (see Figs. 13 & 14, ref 5) and the comparative performance of 628 the various mortars reported here-in since the sand:cement and w/c ratios are different 629 in the two studies. However, the conditioning of the current control mix equates to that of 630 the DARC mortar stored for 4 weeks in [5]. Using these mortars as the basis for 631 comparison the strength of the lime-dried mortar is 134% of the control mortar whilst the 632 strength of the 30 minute stored DARC mortar is 152% of that of the 4 week stored 633 DARC mortar [5]. Thus, the lime-drying has largely mitigated the degradation attributed 634 to the effects of free moisture present during prolonged storage of the DARC mortars; all 635 free moisture would not have been immediately removed by the addition of quicklime so 636 some residual degradation would be expected.

637

Experience suggests that once the water content of the lime-dried mortar is adjusted to yield the desired flow, the strength would be similar to that of the control mix and the workable life extended into the target range of 1 - 2 hours; additional work is required to prove this assumption.

642

643 **7 Conclusions**

644

645 From the results described in Parts A and B of this paper it can be concluded that:

646

The lime-drying process has been successfully employed to produce mortars of M1
 and M5 classification using ggbs as the hydraulic component.

By careful manipulation of the amount of quicklime added (stoichiometric ratio) and
 the length of the mixing time during the lime-drying process it is possible to produce
 identical mortars from sands with differing water contents.

- The composition of different quicklimes may have a significant influence on the
 strength of hardened mortars prepared from the lime drying process, particularly in
 lime-rich blends with low ggbs/CH.
- The strength of the lime-slag mortars produced by the lime-drying process are very similar to those of mortars produced using equivalent commercial hydrated lime.
- The lime dried materials may be silo-stored prior to use without changes to the mortar classification.
- The work has highlighted a requirement for further research to examine the effects of
 silo storage on the subsequent water demand and strength of lime mortars
 containing a high proportion of lime.
- Lime-drying overcomes strength degradation previously observed in Roman cement
 mortars retarded by a pre-hydration technique and subsequently stored.
- 664

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666

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