Rehydroxylation (RHX) dating: Mass loss issues due to incomplete drying, carbon content, and mineral alteration


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Abstract:

As part of rehydroxylation (RHX) dating trials on post-medieval bricks, investigations were carried out into the presence, and effect on age estimations, of uncertainties associated with (a) prolonged and incomplete drying of samples, (b) the removal of organic and non-organic matter during heating, and (c) mineral alteration during heating. All samples exhibit an issue with a prolonged period of drying, exceeding two months at 130°C. Methods for treating this issue are applied and demonstrate the need for higher resolution and more precise monitoring of the mass loss during drying if the moisture not removed needs to be taken account of; this may not be necessary if prolonged drying is associated with the slow removal of chemisorbed water, distinct from rehydroxylation-related water loss. Organic matter contamination is also present in significant quantities in all samples, regardless of the retrieval context; the considerable effects of uncertainties in this quantity, arising from variability in the organic matter to organic carbon ratio (OM/OC), on age estimations are presented. Mineral alteration during reheating is generally negligible except when gypsum is present; large sources of uncertainty arising from moisture loss associated with dehydration and with subsequently lower levels of physisorption following reheating are highlighted as problematic.

Keywords:

Rehydroxylation, Dating, Drying, Contaminants, Minerals, Components
1. **Introduction**

The potential of rehydroxylation dating as an archaeological tool showed early promise (Wilson 2009; 2012) but has since suffered from an absence of successful applications (e.g. Burakov et al. 2013; Le Goff and Gallet 2015a; Numrich et al. 2015) and the identification of a range of issues surrounding the method (e.g. Bowen et al. 2011; 2013; Le Goff and Gallet 2015a; 2015b; Gallet and Le Goff 2015; see also the review in Barrett 2015). These often focus on the mass gain behavior that is central to the method, for example the suitability of a $t^{1/4}$ model (e.g. Bowen et al. 2011; Le Goff and Gallet 2014; 2015b; Gallet and Le Goff 2015; Barrett 2017b) or the equilibration of the mass following drying (e.g. Le Goff and Gallet 2014; Gallet and Le Goff 2015). However, there are also issues surrounding estimation of the amount of mass, due to rehydroxylation (uptake of atmospheric moisture that is subsequently chemically bound to the ceramic matrix), gained by the ceramic since it was last dehydroxylated (i.e. originally fired or last heated above 500°C). This mass is critical to age calculations and can be retrieved based on the mass loss that occurs during reheating at 500°C (Wilson 2012). There may be components of this mass loss that have other origins and need to be taken account of: incomplete drying (Bowen et al. 2011; Barrett 2015) may result in a quantity of loose or capillary water later removed at the higher reheating temperature; organic carbon may be burnt off between 200-500°C (Numrich et al. 2014); and minerals may undergo alteration and contribute additional mass loss, i.e. goethite (Burakov and Nachasova 2013). These mass loss issues, particularly incomplete drying and organic matter, have not received enough focused attention. As part of a larger work aimed at assessment of the archaeological application of RHX dating (Barrett 2015; 2017a; 2017b), these factors were explored, pre-planned in the case of organic contaminants and mineral alteration but necessitated unexpectedly in the case of indefinite drying; the dating trial results are presented in the companion article (Barrett 2017a).

The results of mass loss data for drying granulated samples will be presented together with modelling attempts to account for any remaining moisture (mass) in the fabric. The potential effect this estimated remaining mass has on age estimations from dating trials (Barrett 2015; 2017a) will be explored. As well as this, an explanation for prolonged drying will be suggested.

The presence and nature of organic matter, lost during heating, are examined using Fourier transform infrared spectroscopy (FTIR), carbon content analysis, and thermogravimetric mass spectrometry (TG-MS). The mass loss
uncertainties (and their age estimation impact) that arise from issues with the organic matter to organic carbon (OM/OC) ratio will be explored.

Finally, the effects of reheating on the mineral composition are examined through x-ray diffractometry (XRD) and FTIR.

Note that in the associated dating trials both mass gain models, \( t^{1/4} \) and \( t^{1/n} \), were applied for reasons discussed in Barrett (2017a). While the latter was preferred, the age estimations produced using both approaches were problematic. The present work discusses mass loss effects in relation to both, with preference often given to the use of the \( t^{1/4} \) model results in illustrating the magnitude of potential effects (of relevance in re-examining successful RHX dating trials that used this model, i.e. Wilson et al. 2009; 2012); counterpart effects for the \( t^{1/n} \) model can be found in Barrett (2015) and are not included here if the magnitude of the effect is very similar to that for the \( t^{1/4} \) model such that it does not alter or contribute significantly to the main conclusions and arguments being made.

2. Method

2.1 Samples

The sample set (see supplementary materials Table S1 and S2) is described in more detail (including compositional, surface area, firing information, origin, context) in the companion article (Barrett 2017a) and more comprehensively elsewhere (Barrett 2015). It was composed of 18 samples, including 14 brick samples from the island of Ireland, 2 pottery samples from Northern Ireland, and a piece of Roman and Etruscan ceramics; the brick and pottery samples were of post-medi eval age (Table S1). Samples were retrieved from both buried (9) and non-buried (9) contexts (Table S2). Brick samples (of particular importance as they provided all 12 samples suitable for age estimates in Barrett 2015; 2017a) were likely produced from residual/boulder clays (rich in quartz and feldspars and generally low in calcium content) that were well fired above 850°C (Barrett 2015; 2017a). The specific surface area of samples was quite variable but typically < 5m²/g (Table S.3).

2.2 Drying and Loose Water Modelling

Drying at 130°C was conducted during dating trials (Barrett 2015; 2017a) to remove any non-rehydroxyl moisture from the ceramic (i.e. pore or capillary bulk moisture). Samples had already undergone extensive pre-drying at approximately 80°C for 15 days during preparation before this stage (see Barrett 2015). For drying, granulated
samples (typically > 80g of 2-5.6mm sieved fractions, each granule approx. 0.05-0.15g) were spread out on aluminium trays and placed in an oven (fan assisted) at 130°C (granulation was conducted to homogenize sample material, originally quite heterogeneous, before it was split into three subsamples for dating trials). Based on the existing literature (see Discussion), it was assumed that samples would dry to constant mass within a reasonable period of time (maximum of a couple of weeks). On a near daily basis, the trays were removed and weighed on a top-loading balance under room conditions of 20 ± 1°C and 78 ± 3%RH. Samples were weighed under a rigid timing regime for two months; it was recognized from the mass loss curves that after a month of repeated measurements equilibration to constant mass appeared unlikely with measurements continued for an additional month to improve the quality of the mass loss curves.

Modelling of the drying curves was carried out to provide an estimate of the magnitude of loosely bound water, \( m_{lw} \), not removed during drying (presumed to be removed when samples are reheated at 500°C as part of dating trials). Two empirical models of drying were applied using the curve fitting tool cftool in MATLAB R2012a:

Model 1 (Power):  
\[ m(t) = a(t^{-b})+c \]  
where \( a \) is the mass rate, \( b \) is the power of the model, and \( c \) is the final dry mass of the ceramic.

Model 2 (Exponential):  
\[ m(t) = ae^{-bt} + c \]  
where \( a \) and \( b \) (drying constant) are coefficients that determine the rate of moisture loss and \( c \) the final dry mass of the ceramic.

The former of these models was trialled based on the emphasis of the prevalence of \( t^{1/2} \) transport processes in ceramic materials (Brosnan and Robinson 2003). However, it was found that the \( 1/2 \) power was a poor descriptor of the behaviour observed; hence the power constraint was relaxed.

The latter of these two models is derived from Lewis’ (1921) thin-layer equation:

\[ \frac{dX}{dt} = b(X - X_e) \]  
where \( X \) is the moisture content, \( X_e \) is the equilibrium moisture content (i.e. 0 for an idealised dry ceramic) and \( b \) is the drying constant. Lewis (1921) considered that during drying of porous and hygroscopic materials, the rate of loss of moisture is proportional to the difference between the moisture content and the moisture content when the material
is at equilibrium (dry) with the drying environment. This thin-layer equation assumes that the conditions of drying (humidity, temperature) are constant throughout the material during drying. Forms of this equation are commonly used in fundamental modelling of drying (see Pakoswki and Mujumbar 2006 and Marinos-Kouris and Maroulis 2006).

The power model proved unsatisfactory (see Results and Discussion) and the exponential model was instead selected for complete modelling and calculation of the \( m_{lw} \) values. This model was used to estimate the completely dry mass of the sample trays and the mass at 61, 62, 63 and 66 days (when groups of samples were removed for commencement of mass gain measurements), with the difference between these two amounting to the potential remaining moisture that could be removed upon heating at 500°C. Uncertainties at the 95% confidence level were calculated from the upper and lower bounds on the model coefficients estimated in MATLAB.

2.3 Thermogravimetric Mass Spectrometry

TG-MS was carried out using a Netzsch TG 209 F1 Libra thermo-microbalance in series with a Pfeiffer Thermostar mass spectrometer. Approximately 30-40mg of powdered dating sample (<63μm) was used. The samples had previously been dried at 130°C and stored in a desiccated environment for a period of three months. The sample was placed in an Al₂O₃ crucible within the TG 209 and heated from 25°C to 1000°C at a rate of 20°C/min under a constant flow of nitrogen at 50mL/min. Mass spectrometry was carried out for ions of mass number 18 (H₂O), 44 (CO₂) and 64 (SO₂), liberated during the heating process. Sampling of the mass and ion current (mass spectrometry) was carried out for each °C increment in temperature.

Analysis (identification of events and structures) of the mass loss curve, its first derivative, and ion fragment curves was carried out after smoothing (typically a moving average with \( n = 11 \)).

2.4 Carbon Content and Organic Matter Mass

Carbon content analysis was carried out at the \(^{14}\)CHRONO Centre at Queen’s University Belfast. For each dating sample, carbon content analysis was carried out on two subsamples of approximately 0.5g powdered form (<63μm); one sample that had been dried at 130°C and another that had been fired at 500°C (these subsamples came from powdering of remaining (typically 10-20g) homogenised sample material not used in dating). Samples were combusted (in the presence of copper oxide and strips of silver) in quartz-glass tubes at 850°C for 8 hours. Then the
sample was opened into a cryogenic separation line and the CO₂ gas pressure in a specific volume recorded and converted to a mass of carbon using a laboratory conversion factor. The ratio of the mass of carbon to the original mass of the sample was used to calculate the %wt C (carbon).

For each dating sample, this provided two values, the %wt C for the sample dried at 130°C (corresponding to mass \( m_{c,130°} \)) and the %wt C for the sample heated at 500°C (corresponding to mass \( m_{c,500°} \)). To obtain the %wt C removed from a sample during heating between 130-500°C \( (m_c) \) the following equation was used:

\[
m_c = m_{c,130°} \cdot m_{c,500°}
\]

As part of dating trials (Barrett 2015, 2015b), before the age of the ceramic could be calculated an estimate of the organic matter mass, \( m_{om} \), removed during reheating must be subtracted from the total mass loss during reheating. In order to do this the carbon content must be multiplied by an organic matter to organic carbon ratio, \( OM/OC \):

\[
m_{OM} = OM/OC \cdot m_c
\]

For age calculations, a standard value of 1.95 was used for \( OM/OC \) across all samples. For assessment of the effect of variation in the \( OM/OC \), a range of 1.4-2.5 was used (Pribyl 2010; see Barrett 2015 Section 2.5 for a summary review of the literature upon which these values were selected).

### 2.4 FTIR and XRD

FTIR analysis (Russell and Fraser 1994) was carried out using a PerkinElmer Spectrum One. For each dating sample, two powdered subsamples (< 63 μm) were prepared into pressed pellets, a sample that had previously been dried at 130°C (3 months) and a sample that had been fired at 500°C (for 18 hours). This was achieved by mixing approximately 2-5mg of sample with 40-100mg of potassium bromide (KBr) and compressing the pellet in an evacuated die under high pressure (10 tons) to form a pressed disc of diameter 13mm and thickness <1mm. These discs were then dried in an oven at 130°C for a minimum of 3 hours to remove any loosely bound surface adsorbed water before being placed in a desiccated jar to cool. Scans were then run on room temperature samples over the wavenumber range 450cm\(^{-1}\) – 4000cm\(^{-1}\). The Spectrum One software was used to carry out a baseline correction on each of the scans.
Using MATLAB, the second derivative of the FTIR spectrum and a smoothed version of this second derivative (using cftool and a smoothing spline with smoothing parameter 0.0133) were acquired. The main absorption peaks were identified through comparison of the FTIR scan and its $2^{nd}$ derivative with the mineral identification (carried out largely over the region 1500-450cm$^{-1}$) using tables compiled from the following references: Benedetto et al. 2002; Chukanov 2014; Russell and Fraser 1994; van der Marel and Beuterspacher 1976.

Special attention was also paid to the region 3000-2800cm$^{-1}$ where absorption peaks associated with organics were compiled and tabulated.

XRD analysis was also carried out on powdered (<63μm) subsamples of both dried (130°C for 3 months) and reheated (500°C for 18hrs) dating samples. Approximately 1-2 g of powder, enough to fill a 16mm diameter holder of depth 2.4mm, was required per subsample. X-ray diffractometry was then carried out with a PANalytical X’Pert Pro system using Cu-Kα radiation. Scans were run over the range 3-63° (2θ) with a step size of 0.0170 (2θ) and scan step time of 120s (total scan time of 58 minutes per sample).

Analysis of the XRD spectra (Moore and Reynolds 1997) was carried out using the powder pattern analysis tool PANalytical X’Pert Highscore Plus (Highscore 2015). The spectra had the following treatments carried out in the following order: background removal; smoothing (quantic polynomial over a window size of n=15); peak search (using the minimum of the $2^{nd}$ derivative method and a minimum peak significance of 15 counts); convert FDS to ADS correction; search and match algorithm to identify the possible minerals present using the International Center for Diffraction Data (ICDD) library PDF2. Identification of minerals present was based on both peak and profile matching with the library spectra available.

3. Results

3.1 Drying, Loose Water and Effects

The goodness-of-fit results ($R^2$ and RMSE) of the application of both drying models to the drying mass curves are presented in Table 1 with examples of the modelled fits displayed in Figure 1 for both well behaved (Joy, top) and abnormally (Lan, bottom) behaved samples. The terminology ‘well behaved’ and ‘abnormal’ samples corresponds to samples that displayed mass gain curves that were well suited (a clear Stage 2 behaviour that is linear as a function of
together with an Arrhenius temperature dependence for the mass gain rate) or poorly suited for age estimate modelling; this is discussed in the companion article (Barrett 2017a). Differences in the goodness-of-fits between the two drying models are minor and alone are not significant enough to merit giving preference to either model.

To further examine the strength of both models, the modelled loose water content, as a fraction of the modelled dry mass, was correlated against properties of the ceramic that would be expected to have a strong/moderate relationship with moisture content (%wt. loss between 50-130°C from TG-MS results, BET surface area and pore volume (Barrett 2015; 2017a; 2017b, see also Table S3)), Table 2, and provide stronger support for the use of Model 2.

Table 1: Comparison of goodness-of-fit estimates for two models applied to drying curves of dating samples.

<table>
<thead>
<tr>
<th></th>
<th>Model 1 ((m(t)=a(t^b)+c))</th>
<th>Model 2 ((m(t)=a(exp^{-bt})+c))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(R^2)</td>
<td>RMSE (g)</td>
</tr>
<tr>
<td>Ann</td>
<td>0.67</td>
<td>0.008</td>
</tr>
<tr>
<td>Esp</td>
<td>0.16</td>
<td>0.006</td>
</tr>
<tr>
<td>Nic</td>
<td>0.72</td>
<td>0.008</td>
</tr>
<tr>
<td>Mac</td>
<td>0.90</td>
<td>0.009</td>
</tr>
<tr>
<td>Ria</td>
<td>0.96</td>
<td>0.008</td>
</tr>
<tr>
<td>Etr</td>
<td>0.81</td>
<td>0.025</td>
</tr>
<tr>
<td>Rom</td>
<td>0.82</td>
<td>0.016</td>
</tr>
<tr>
<td>Por</td>
<td>0.90</td>
<td>0.017</td>
</tr>
<tr>
<td>Rat</td>
<td>0.89</td>
<td>0.006</td>
</tr>
<tr>
<td>Cal</td>
<td>0.95</td>
<td>0.007</td>
</tr>
<tr>
<td>Lan</td>
<td>0.72</td>
<td>0.016</td>
</tr>
<tr>
<td>Joy</td>
<td>0.96</td>
<td>0.008</td>
</tr>
<tr>
<td>Cau</td>
<td>0.78</td>
<td>0.008</td>
</tr>
<tr>
<td>Bel</td>
<td>0.90</td>
<td>0.013</td>
</tr>
<tr>
<td>Dow1</td>
<td>0.81</td>
<td>0.017</td>
</tr>
<tr>
<td>Dow2</td>
<td>0.91</td>
<td>0.009</td>
</tr>
<tr>
<td>Tur</td>
<td>0.92</td>
<td>0.009</td>
</tr>
<tr>
<td>Ted</td>
<td>0.86</td>
<td>0.008</td>
</tr>
</tbody>
</table>
Figure 1: Drying mass curves for *Joy* (top) and *Lan* (bottom) with two models applied. Red dashed = exponential model; blue dotted = power model.
Table 2: Assessment, for both drying models, of the level of correlation ($R^2$) between the remaining water content (as a fraction of the estimated dry mass) and various proxies for the potential moisture content of the sample (BET S. A. and pore volume from Barrett 2015). From linear regressions carried out across all dating samples.

<table>
<thead>
<tr>
<th>versus</th>
<th>TG-MS 50-130°C %wt loss</th>
<th>BET Surface Area</th>
<th>Pore Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model 1 ($R^2$ value)</td>
<td>0.11</td>
<td>0.04</td>
<td>0.10</td>
</tr>
<tr>
<td>Model 2 ($R^2$ value)</td>
<td>0.53</td>
<td>0.38</td>
<td>0.43</td>
</tr>
</tbody>
</table>

The modelled loose water content remaining in the dried samples, after 60 days drying and given as a percentage of the modelled dry mass, is presented in Figure 2 and arranged in order of dryness with 95% confidence intervals included also. It is notable that many of the samples with the largest values of loose water are samples (Etr, Lan, Dow1/2, Por, Mac,Bel) considered problematic in related work due to having a high specific surface area (Table S.3) that results in potential capillary condensation issues (Barrett 2015; 2017b). The sample Esp is significant as it appears to be the only sample to have dried out.

Figure 2: The modelled loose water content, as a percentage of the modelled dry mass of samples (Model 2), remaining after 60 days drying at 130°C. Arranged in order of dryness and with 95% confidence intervals.

For examination of the implications with regard to RHX dating, the mass of loose water, $m_{lw}$, is presented as a percentage of the total mass loss, $m_{RHX}$, during reheating at 500°C (with regard to application of a $t^{1/4}$ model; similar
results apply to a $t^{1/n}$ model, Figure 3. These values vary, typically, between 0.25-1.5% but with an exceptionally high value of 3.2% (Bel, $t^{1/4}$ model) also obtained.

![Figure 3: The mass of remaining loose water, $m_{lw}$, as a % of total fractional mass loss, $m_{RNA}$, upon heating between 130°C and 500°C. For $t^{1/4}$ model.](image)

### 3.2 TG-MS

The full set of TG-MS curves are provided in Barrett (2015). A summary of the TG-MS features observed are presented in Table 3. To illustrate the main heating events in this table associated with CO$_2$ (organic/inorganic carbon) loss, Figure 4 presents the mass spectrometry curve for mass 44 (CO$_2$) with events E-G labelled. The CO$_2$ associated events occur over the temperature regime 200-800°C and are classified as follows:

- **E:** 200-300°C – onset of organic CO$_2$ mass loss.
- **F:** 400-500°C – organic CO$_2$ peak (‘increasing’ implies peak not clear due to overlap with inorganic peak.
- **G:** 600-750°C – inorganic CO$_2$ peak (carbonates).

<table>
<thead>
<tr>
<th>Event</th>
<th>Temperature Range</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>E</td>
<td>200-300°C</td>
<td>Organic CO$_2$ mass loss onset</td>
</tr>
<tr>
<td>F</td>
<td>400-500°C</td>
<td>Organic CO$_2$ peak (‘increasing’ implies peak not clear due to overlap with inorganic peak)</td>
</tr>
<tr>
<td>G</td>
<td>600-750°C</td>
<td>Inorganic CO$_2$ peak (carbonates)</td>
</tr>
</tbody>
</table>

Table 3: Summary of TG-MS analysis. Significant peaks (corresponding to events A-D) where weight loss is attributed to H$_2$O loss are colour coded according to strength of visible structure (all numbers are in °C). Onset of organic CO$_2$ mass loss (event E) and organic CO$_2$ maximum (event F) are included (‘increasing’ implies no
clear peak and overlap with inorganic calcite-related peak) together with position of inorganic carbon (calcite-related) peak (event G). The presence/absence of SO\textsubscript{2} removal is also provided.

<table>
<thead>
<tr>
<th>Event</th>
<th>H\textsubscript{2}O peaks (yellow = weak, orange = medium, red = strong)</th>
<th>CO\textsubscript{2} onset</th>
<th>Organic CO\textsubscript{2} max.</th>
<th>Inorganic CO\textsubscript{2} max.</th>
<th>SO\textsubscript{2} (a=absent; p=present )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ann</td>
<td>70</td>
<td>-</td>
<td>250-300</td>
<td>450-500</td>
<td>620</td>
</tr>
<tr>
<td>Esp</td>
<td>60-70</td>
<td>110</td>
<td>260-270</td>
<td>-</td>
<td>250-300</td>
</tr>
<tr>
<td>Nic</td>
<td>50</td>
<td>- 210-230</td>
<td>- 250-300</td>
<td>increasing 610-620</td>
<td>400</td>
</tr>
<tr>
<td>Mac</td>
<td>90</td>
<td>- 310-320</td>
<td>200-250</td>
<td>410-420</td>
<td>trace</td>
</tr>
<tr>
<td>Ria</td>
<td>50</td>
<td>120-160</td>
<td>- 250-300</td>
<td>450-500</td>
<td>610-620</td>
</tr>
<tr>
<td>Etr</td>
<td>90</td>
<td>240</td>
<td>- 250-300</td>
<td>450-500</td>
<td>740-750</td>
</tr>
<tr>
<td>Rom</td>
<td>90-100</td>
<td>- 230-240</td>
<td>- 250-300</td>
<td>450-500</td>
<td>710-720</td>
</tr>
<tr>
<td>Por</td>
<td>60</td>
<td>160-170</td>
<td>- 350</td>
<td>250-300</td>
<td>increasing 600-610</td>
</tr>
<tr>
<td>Rat</td>
<td>60</td>
<td>140</td>
<td>- 340-350</td>
<td>300-350</td>
<td>430-450</td>
</tr>
<tr>
<td>Cal</td>
<td>90</td>
<td>- 210</td>
<td>300-350</td>
<td>430-450</td>
<td>630-640</td>
</tr>
<tr>
<td>Lan</td>
<td>90</td>
<td>- 310-320</td>
<td>300-350</td>
<td>430-450</td>
<td>640-650</td>
</tr>
<tr>
<td>Joy</td>
<td>60</td>
<td>150</td>
<td>- 250-300</td>
<td>450-500</td>
<td>640-650</td>
</tr>
<tr>
<td>Cau</td>
<td>70</td>
<td>130-140</td>
<td>270-280</td>
<td>250-300</td>
<td>450-500</td>
</tr>
<tr>
<td>Bel</td>
<td>90-100</td>
<td>- 250-300</td>
<td>230-250</td>
<td>480</td>
<td>680</td>
</tr>
<tr>
<td>Dow1</td>
<td>80-90</td>
<td>- 340-390</td>
<td>250-300</td>
<td>370</td>
<td>730-740</td>
</tr>
<tr>
<td>Dow2</td>
<td>95-110</td>
<td>-</td>
<td>250-300</td>
<td>480-530</td>
<td>750</td>
</tr>
<tr>
<td>Tur</td>
<td>60-70</td>
<td>140-160</td>
<td>250-300</td>
<td>450-500</td>
<td>570-580</td>
</tr>
<tr>
<td>Ted</td>
<td>-</td>
<td>170-180</td>
<td>270-300</td>
<td>450</td>
<td>650-670</td>
</tr>
</tbody>
</table>

All samples featured loss of organic carbon. No inorganic carbon peak was observed for Mac. The presence of SO\textsubscript{2} was observed for Mac and Bel with SO\textsubscript{2} removal occurring above 650-700°C.

Four typical examples of the mass loss due to CO\textsubscript{2} (organic and inorganic) are presented in Figure 5. Clear onsets for CO\textsubscript{2} loss occur in the region 250-350°C, typically, with a plateau of emission common from 450°C until mass loss associated with carbonates commences from 650-700°C (the association of this peak with carbonate minerals is due to their presence in XRD, FTIR and petrography and is expanded upon in Barrett 2015). The curve for Mac (top right) is included as it demonstrates only a trace amount of CO\textsubscript{2} associated with carbonate breakdown.
Figure 4: TG mass loss curve (red) and mass 44 (CO₂) mass spectrometry curve (green) for Esp. Events E-G (see text) are highlighted.

Figure 5: TG mass loss curves (red) and mass 44 (CO₂) mass spectrometry curves (green) for Ann (top left), Mac, (top right), Cal (bottom left), Cau (bottom right).
3.3 Carbon Content

The results of carbon content analysis are provided in Figure 6, presented as %wt C. It can be observed that the magnitude and presence of carbon is not related to whether or not the sample was retrieved from a buried context.

The mass of organic matter (computed using $m_c$ and an OM/OC ratio of 1.95), $m_{om}$, as a % of the total fractional mass loss, $m_{RHXC}$, of dating trial samples (Barrett 2015;2017a) is presented in Figure 7 (for $t^{1/4}$ model, similar levels for $t^{1/n}$, see Figure 5 of Barrett 2017a). Levels of 5-20% are common with some values also in the range 35-45%.

The effect uncertainties in the OM/OC multiplicative factor (1.4-2.5) have on the age estimates from dating trials are displayed in Table 4, for both the $t^{1/4}$ and $t^{1/n}$ models. For the examples Joy and Ann, age-temperature curves, which display the age estimations as a function of effective lifetime temperature (ELT), are provided in Figure 8 (see Barrett 2017a for details and Barrett 2015 for full set of curves).

Figure 6: Percentage weight of carbon removed from dating samples during firing at 130-500°C. The retrieval of the sample from a buried context is indicated by (b).
Table 4: Examples of the effect of uncertainties in OM/OC ratio (using 2.5 or 1.4) on the age range estimates.
For $t^{1/4}$ and $t^{1/n}$ model. Italicised and bold correspond to age ranges that overlap with known age.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Known Age (Yrs)</th>
<th>OM/OC (2.5)</th>
<th>OM/OC (1.4)</th>
<th>OM/OC (2.5)</th>
<th>OM/OC (1.4)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ann</td>
<td>110±14</td>
<td>319</td>
<td>1283</td>
<td>59</td>
<td>182</td>
</tr>
<tr>
<td>Esp</td>
<td>141±6</td>
<td><em>147</em></td>
<td><em>452</em></td>
<td>284</td>
<td>1115</td>
</tr>
<tr>
<td>Nic</td>
<td>398±2</td>
<td>46440</td>
<td>66885</td>
<td>820437</td>
<td>1267480</td>
</tr>
<tr>
<td>Mac</td>
<td>228±2</td>
<td>477321093</td>
<td>642239904</td>
<td>8113299</td>
<td>10271317</td>
</tr>
<tr>
<td>Ria</td>
<td>339±25</td>
<td>122179</td>
<td>191242</td>
<td>4150</td>
<td>5660</td>
</tr>
<tr>
<td>Rat</td>
<td>245±2</td>
<td>1415987</td>
<td>2145539</td>
<td>4343</td>
<td>5517</td>
</tr>
<tr>
<td>Cal</td>
<td>182±9</td>
<td>2337</td>
<td>3014</td>
<td><em>177</em></td>
<td><em>212</em></td>
</tr>
<tr>
<td>Joy</td>
<td>412±2</td>
<td>321</td>
<td>489</td>
<td>152</td>
<td>208</td>
</tr>
<tr>
<td>Cau</td>
<td>399±4</td>
<td><em>388</em></td>
<td><em>704</em></td>
<td>--</td>
<td>--</td>
</tr>
<tr>
<td>Bel</td>
<td>395±3</td>
<td>258</td>
<td>2178</td>
<td>774</td>
<td>17058</td>
</tr>
<tr>
<td>Tur</td>
<td>229±35</td>
<td>36508</td>
<td>62139</td>
<td>359</td>
<td>494</td>
</tr>
<tr>
<td>Ted</td>
<td>339±25</td>
<td>174</td>
<td>263</td>
<td>16</td>
<td>21</td>
</tr>
</tbody>
</table>

Figure 7: The mass of organic matter (computed using $m_c$ and an OM/OC ratio of 1.95), $m_{om}$, as a % of total fractional mass loss, $m_{RHXC}$, upon heating between 130°C and 500°C. For $t^{1/4}$ model.
Figure 8: Effect of uncertainties in OM/OC ratio (1.4-2.5, green dash) on the age temperature curves of Joy (top) and Ann (bottom). Red line is known age. Black dashed line is ELT estimate. Using a $t^{1/4}$ model.
3.4 FTIR/XRD

3.4.1 Reheating Mineral Alteration

The complete collection of spectra with tables of peak and mineral identification for all samples using both XRD and FTIR are provided in Barrett (2015). With the exception of two samples, Mac and Bel, the spectra from XRD and FTIR showed no signs of mineral alteration associated with reheating, see Figure 9 for Ann for the typical level of similarity in the reheated and non-reheated spectra.

![Figure 9: XRD (top) and FTIR (bottom) Spectra of non-reheated and reheated Ann sample. Inset of the region 450-900 cm⁻¹ used in mineral identification.](image-url)
Two samples, *Mac* and *Bel*, displayed significant differences in spectra, with *Figure 10 (top)* for *Mac* representative of the major XRD differences observed in *Bel* also. For *Mac*, the XRD spectrum of non-reheated sample has peaks at 14.7°, 29.7° and 31.8° (2θ) that are either non-existent or very weak in the reheated sample spectrum, whereas the reheated sample spectrum has peaks at 25.4°, 38.6°, 48.6° and 52.3° (2θ) that are either non-existent or significantly weaker in the non-reheated sample. The former non-reheated peaks have been identified as bassanite (hemihydrate) reflections and the latter reflections in the reheated samples are attributed to anhydrite (using ICDD PDF2 peak identification library). Differences in the two spectra are attributed to the level of hydration of sulfates within the ceramic. For *Bel*, there are strong differences in the non-reheated and reheated spectra. The following peaks are present in the non-reheated and either absent or very low-level in the reheated sample: 14.6°, 29.6°, 31.8°(2θ). These can all be identified with bassanite (hemihydrate). For the reheated sample, peaks not present in the non-reheated sample are: 24.1°(2θ), 25.4°(2θ), 31.3°(2θ), 38.6°(2θ), 40.8°(2θ) and 41.6°(2θ). These can all be identified with anhydrite. The differences are similar to those recorded for *Mac*.

This is supported by the FTIR spectra, shown for *Mac* in *Figure 10 (bottom)* and representative of the differences observed for *Bel*. There are some significant differences between the non-reheated and reheated *Mac* samples. The reheated sample has peaks at 575cm⁻¹ and 614cm⁻¹ that are either very weak or not present in the non-reheated samples as well as a peak at 677cm⁻¹ that appears to be greater in magnitude and shifted from 673cm⁻¹ in the non-reheated sample. The non-reheated sample has a peak at 633cm⁻¹ that is not present in the reheated sample together with the previously mentioned shifted peak at 673cm⁻¹. The following minerals can be attributed to the aforementioned peaks: Anhydrite = 575, 614, 617, 673, 677cm⁻¹; Bassanite = 633cm⁻¹. The change upon reheated is associated with dehydration of sulfates (gypsum).

For *Bel*, peaks at 663cm⁻¹, 669cm⁻¹ and 1154cm⁻¹ in the non-reheated samples are consistent with bassanite, gypsum and bassanite, respectively. Peaks at 613cm⁻¹ and 678cm⁻¹ in the reheated sample are attributed to anhydrite. There are no other significant differences between the spectra aside from those associated with variations in adsorbed moisture and atmospheric carbon dioxide.
Figure 10: Top - XRD spectra of Mac sample without (red) and with (blue) reheating at 500°C. Region constrained to 10-55° (2θ). Red ovals mark basanite (hemihydrate) peaks; blue ovals mark anhydrite peaks. Bottom - FTIR spectra of non-reheated (blue) and reheated (red) Mac sample over the region 450-900(1/cm). Gypsum associated peaks (anhydrite/bassanite) are highlighted together with some examples of quartz and feldspars.
3.4.2 TG-MS Gypsum

The TG-MS measurements provided evidence for release of SO$_2$ only for samples Mac and Bel, see Figure 11 for Bel, and with an onset above 650-700°C.

![Figure 11: TG mass loss curve (red) and mass 64 (SO$_2$) mass spectrometry curve (black) for Bel.](image)

3.4.3 Stage 1 Behaviour

Dealt with in Barrett (2015; 2017b) but of relevance to the present work are the Stage 1 (S1) portion of mass gain curves obtained following drying (130°C) and reheating (500°C) of samples. Stage 1 corresponds to the initial and more rapid mass gain that occurs following drying/reheating and is attributed to physisorption-related processes (Barrett 2015; 2017b); Stage 1 commences with exposure of the ceramic to moisture in the air and is considered complete with the commencement of Stage 2, i.e. when the mass gain curve is adequately modelled by a $t^{1/6}$ or $t^{1/4}$ model alone. For most samples, the magnitude of mass gain in this stage was very similar following both drying, $m_{S1,130}$, and reheating, $m_{S1,500}$; however, for Mac and Bel, there were considerable differences, see Figure 12 for Mac. The magnitudes (fractional) are provided in Table 5 with a difference of the order of ~ 4 evident.
Table 5: Stage 1 fractional mass for 130C and 500C components (25°C aging) together with the ratio of the two. For the $t^{1/4}$ model (see Barrett 2015 for the method).

<table>
<thead>
<tr>
<th>$t^{1/4}$</th>
<th>$m_{S1-130}$</th>
<th>$m_{S1-500}$</th>
<th>$m_{S1-130} / m_{S1-500}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mac</td>
<td>0.0030745</td>
<td>0.0007159</td>
<td>4.30</td>
</tr>
<tr>
<td>Bel</td>
<td>0.0048373</td>
<td>0.0012858</td>
<td>3.76</td>
</tr>
</tbody>
</table>

Figure 12: Fractional mass gain curves, displaying the Stage 1 region, of Mac sample. Sample aging at 25°C following heating at 130°C (blue) and 500°C (red). Normalised relative to the Stage 2 modelled intercept mass for display purposes.
3.4.4 Organics FTIR

The results of analysis of the region 2800-3000cm\(^{-1}\) for organics are presented in Table 6, and Figures 13-15. Table 6 presents, for all samples and for reheated and non-reheated spectra, the peak positions (cm\(^{-1}\)) and a subjective assessment of the strength of those peaks. It also attempts to group the observed peaks. Figure 13 is a typical example of the spectra in the region of interest, 2800-3000cm\(^{-1}\). Figure 14 presents the peak position and their possible ranges as a function of wavenumber for non-reheated and reheated samples. Using the peak position ranges in Figure 14, the number of samples with organic peaks at wavenumbers from 2840-2980cm\(^{-1}\) is presented in Figure 15. This data is smoothed with a running average of n = 9 (cm\(^{-1}\)).

![Figure 13: FTIR spectra across region of interest in organic identification for Mac (non-reheated and reheated). Peaks in the ranges 2850-2860cm\(^{-1}\), 2920-2930cm\(^{-1}\), 2950-2970cm\(^{-1}\) are typical for most samples.](image-url)
Table 6: FTIR peaks positions in organics regions 2800-3000 cm\(^{-1}\) for dating samples, both non-reheated and reheated. Colour denotes relative strength of peaks: green=weak; gold=moderate; red=strong. Average values and standard deviations are based on midpoints of ranges.

<table>
<thead>
<tr>
<th></th>
<th>Non-Reheated</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Ann</td>
<td>Esp</td>
<td>Nic</td>
<td>Mac</td>
<td>Ria</td>
<td>Etr</td>
<td>Rom</td>
<td>Por</td>
<td>Rat</td>
<td>Cal</td>
<td>Lan</td>
<td>Joy</td>
<td>Cau</td>
<td>Bel</td>
<td>Dow1</td>
<td>Dow2</td>
<td>Tur</td>
<td>Ted</td>
<td>Avg</td>
<td>Std. Dev.</td>
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<td></td>
<td>2866</td>
<td>2855</td>
<td>2850-2870</td>
<td>2873</td>
<td>2850</td>
<td>2850-2890</td>
<td>2856</td>
<td>2855-2885</td>
<td>2855-2855</td>
<td>2850-2855</td>
<td>2850-2855</td>
<td>2850-2855</td>
<td>2850-2890</td>
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<td>2930-2970</td>
<td>2922</td>
<td>2923-2960</td>
<td>2922-2969</td>
<td>2925-2930</td>
<td>2925-2930</td>
<td>2920-2925</td>
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<td>2960</td>
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</tbody>
</table>

Figure 15: Moving average (n=9) of number of samples with organics peaks at wavenumbers from 2840-2980 cm\(^{-1}\), for both non-reheated (blue) and reheated (red).
4. Discussion

4.1 Incomplete Drying

Two models (power-based and exponential-based) were applied to the drying mass loss curves, Figure 1, with the quality of the regressions quite variable, Table 1. Differences between the goodness-of-fits of both models are minor and do not provide clear support for the selection of either. A better distinction might be made with improved quality data; however, this issue with prolonged drying was unexpected when the experiment was designed. Control of environmental conditions during removal from the oven for weighing was inadequate to prevent large fluctuations in the mass, associated with the level of physisorption taking place; this is reflected in the RMSE values of the models, Table 1, where samples with high surface area (Etr, Rom, Por, Lan, see Table S.3) have the largest fluctuations, see for example Figure 1 for Lan.

The correlation of the modelled remaining mass of loose water with properties with which it would be expected to be associated, for example surface area and pore volume, was examined, Table 2; these results favour the exponential model.

The exponential model was ultimately selected for the reasons above and also because the power-based model was problematic; for a number of samples, the modelled mass of remaining loose water exceeded the mass loss estimates due to heating at 500°C, and for several samples regressions using the power-based model produced parameter values that were nonsensical (i.e. a final ‘dry’ mass that exceeded the initial mass of the drying curve). As well as this, the use of a power model was motivated by a $t^{1/2}$ behaviour suggested by Brosnan and Robinson (2003); however, the model could only be successfully applied with the power constraint relaxed ($1/t$ instead). The exponential model is more satisfactory in its interpretation, with the rate of moisture removal being proportional to the level of moisture remaining.

While the exponential model was selected for use, the authors emphasises that this was borne out of pragmatism and the appropriateness of this model clearly requires more investigation; the quality of the fits produced, Figure 1, were less than satisfactory, and the confidence intervals were also very large, for example Figure 2. Nonetheless, its use was deemed the only way to proceed. That the modelled remaining loose water was generally greatest for the high surface area samples, Figure 2 and see Table S.3, is somewhat reassuring and would also explain the greater correlation
of this model with the surface area, pore volume and TG-MS data, *Table 2*, the inference being that for high surface area samples a significant quantity of physisorbed water is possibly still being removed when heated at 500°C.

*Table 7: Summary of the drying conditions and presence of drying data in the literature.*

<table>
<thead>
<tr>
<th>Publication</th>
<th>Sample Size</th>
<th>Drying Temp. (°C)</th>
<th>Duration of Drying</th>
<th>Drying Data Presented (Y/N) (other drying proxy data)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wilson et al. (2009)</td>
<td>3-5g</td>
<td>105</td>
<td>Not specified (to constant mass)</td>
<td>N</td>
</tr>
<tr>
<td>Bowen et al. (2011)</td>
<td>32-96g (sherd)</td>
<td>110</td>
<td>20-30hrs (However, incomplete drying after 2 weeks)</td>
<td>Y</td>
</tr>
<tr>
<td>Wilson et al. (2012)</td>
<td>0.5-4.0g</td>
<td>105</td>
<td>To constant mass (‘few hours’ to ‘several days’)</td>
<td>N</td>
</tr>
<tr>
<td>Bowen et al. 2013</td>
<td>1-2g</td>
<td>110</td>
<td>30-50hrs</td>
<td>N</td>
</tr>
<tr>
<td>Burakov and Nachasova (2013)</td>
<td>1-15g</td>
<td>105</td>
<td>12hrs</td>
<td>N</td>
</tr>
<tr>
<td>Le Goff and Gallet (2014)</td>
<td>2-3g</td>
<td>105</td>
<td>4 and 16hrs for one group 66 and 51hrs for one group Issue with duration of raised, additional mass loss with repeated heating</td>
<td>N (Mass gain curves demonstrating variation in mass level with drying temp. 95/105°C and duration)</td>
</tr>
<tr>
<td>Le Goff and Gallet (2015a)</td>
<td>2-3g</td>
<td>105</td>
<td>11hrs and 14 days</td>
<td>N (Mass gain curves demonstrating increase in mass gain rate (t^1/2) with prolonged drying)</td>
</tr>
<tr>
<td>Numrich et al. (2015)</td>
<td>2-4g</td>
<td>110</td>
<td>48hrs</td>
<td>N</td>
</tr>
<tr>
<td>Gallet and Le Goff (2015)</td>
<td>1.46-3g</td>
<td>60, 85, 105, 130</td>
<td>5hrs at 60, 85, 105, 130°C 23hrs at 105°C 246hrs at 105°C</td>
<td>N (Mass gain curves demonstrating extra loss in mass as a function of heating temperature and additional heating duration, with effects on mass gain behaviour also evident)</td>
</tr>
<tr>
<td>Zhao et al. 2015</td>
<td>0.4-1g</td>
<td>110</td>
<td>5-7 days at 110°C Overnight at 300°C</td>
<td>N</td>
</tr>
</tbody>
</table>
For all samples, excluding Esp, drying was incomplete after 60 days, Figure 2, with the potential for considerable quantities of moisture to be removed with additional drying, typically 0.25-1.5% of the total mass lost during reheating at 500°C, Figure 3. This poses potential issues for dating estimations where it is necessary to estimate the rehydroxyl mass loss, $m_{RHX}$, during heating at 500°C. If, say, up to 2% of this mass estimate is actually loosely bound physisorbed water, $m_{lw}$, that has not been recognized or poorly accounted for, then a discrepancy in the age estimate of approximately 8% of the age estimate can result (i.e. if there is an uncertainty, $dm$, in the mass, $m$, then the uncertainty, $dt$, in the age estimate, $t$, is given by $(t+dt)/t = ((m+dm)/m)^4$). For the associated dating trial (Barrett 2017a), the estimated $m_{lw}$ was taken account of using the results presented in the present work. There are clearly large uncertainties in this estimation, on account of the quality of the data and uncertainty in the model used. Nonetheless, even permitting $m_{lw}$ to contribute a maximum of 4% of the total mass loss, the result is only a 17% discrepancy in the age estimate and this, unfortunately, is still only a minor contributor to the significant discrepancies observed in age estimates (Barrett 2017a, see also Table 4). Ignoring the results of the dating trial, if all other issues could be removed, then uncertainties of this magnitude alone would be problematic for the validation and use the method.

Prolonged drying is clearly an issue for RHX dating; its possible presence in previous work was examined, with the conditions of drying and evidence for constant mass or long drying times summarised in Table 7. For the majority of these studies, samples were dried either at 105-110°C until they reached constant mass (Wilson et al. 2009; 2012) or for a fixed duration expected to be long enough to dry them out, yet without a strong rational for selection of drying times that range from a few hours to a few days (Bowen et al. 2013; Burakov and Nachasova 2013; Numrich et al. 2015). These studies used samples that are of typical size 0.5-10g, greater than the size of the individual granules used in the present work (maximum size of approx. 0.3g). For experiments where a fixed duration was used, without drying to constant mass being recorded, there is no certainty the samples actually did dry out. For the experiments where samples were stated as drying to constant mass (Wilson et al. 2009; 2012), no evidence or data to support this is presented.

There is support for the findings of the present work in the publications of Bowen et al. (2011), Le Goff and Gallet (2014; 2015a), Gallet and Le Goff (2015) and Zhao et al. (2015). Bowen et al. (2011, Figure 3) produced drying curves that demonstrated that after a period of 2 weeks drying at 100°C constant mass had still not been achieved; however, they do not pursue the matter much further and this was for relatively large samples, 32-96g. Le Goff and
Gallet (2014; 2015a) draw attention to the need for better understanding of the effects of drying temperature and
duration, finding that for longer drying times or higher temperatures there is an additional/greater mass loss
(observable in a shift in the subsequent mass gain curves to lower mass, see Figure B of Le Goff and Gallet (2014)).
Unfortunately, they do not record the mass of the samples as a function of drying time, so again it is not possible to
know if, even for the longest drying times used, 14 days, complete drying occurred. From Gallet and Le Goff (2015)
there is strong evidence that incomplete drying occurs for the duration used (5hrs) while heating at lower temperatures
(60, 85, 105, 130°C) with increases to the next highest temperature producing greater mass loss and mass gain rates.
In Zhao et al. (2015) issues with incomplete drying at 110-120°C in unpublished work is stated as prompting the use
of a drying temperature of 300°C.
In any case, from the above, there is a lack of evidence in the literature that drying to constant mass is actually
achieved, with evidence that it is instead prolonged (Bowen et al. 2011, Zhao et al. 2015) and poorly understood (Le
Goff and Gallet (2014; 2015a); Gallet and Le Goff 2015). The present work suggests that drying is a non-trivial issue
such that even after two months drying at temperatures of 130°C and granules of < 0.2g there is still considerable
potential for water to be removed.
As well as this, there is the question over the nature of the water being removed; is it solely physisorbed/pore/capillary
water that is loosely held/bound (Wilson et al. 2009; 2012) or is chemisorbed water also being removed (e.g. Gallet
and Le Goff 2015; Barrett 2015; 2017b), with the latter responsible for the prolonged behaviour, particularly in low
surface area well-behaved samples (see Barrett 2015; 2017b). If the remaining moisture not removed after 60 days
drying is attributed to loose water, \( m_{lh} \), then it needs to be subtracted from the total mass loss, \( m_{RHXC} \), upon reheating
at 500°C in order to obtain the RHX related mass loss, \( m_{RHX} \) (Barrett 2015; 2017a). However, this assumption of
drying only removing physisorbed water was originally guided by assertions that equilibration of the mass occurred
following drying (Wilson et al. 2009; 2012). The work of Le Goff and Gallet (2014; 2015a), Gallet and Le Goff
(2015) and Barrett (2015) demonstrate that this is not the case, and, further, that those assertions were not well
supported experimentally (Le Goff and Gallet 2015a). Instead, it seems increasingly likely (Gallet and Le Goff 2015;
Barrett 2015; 2017b), that the two stage mass gain behavior, with Stage 2 described well by a \( t^{1/n} \) model, following
drying is a chemisorption driven process and associated with the removal of chemisorbed water during drying.
Interestingly, this seems supported by the result that the only sample to fully dry was Esp for which Event B (TG-MS
data, Table 3, associated with a chemisorption process by Barrett 2015; 2017b) was low in magnitude and with a peak water removal temperature of 110°C, the lowest for samples where Event B was certain, and well below the drying temperature of 130°C used in mass gain tests.

If prolonged drying is associated with chemisorption (with removal of loose water completed at some earlier point) then it may not be necessary to estimate or subtract the quantity $m_{lw}$ if the component based approach proposed by Barrett (2015; 2017) is valid. This approach will not be discussed but provided all physisorbed water is removed then the remaining chemisorbed water not removed is taken account of in its methodology. The implications of this are dealt with in Barrett (2015).

In any case, the present work demonstrates that complete drying is unlikely or impractical using simple oven based setups and highlights the need for more controlled recording of the mass loss curves during drying in order to (a) assess the quantity of loose water remaining (if drying is associated with loose water removal alone) or (b) to provide evidence that the stage associated with removal of loose water is complete, with continued mass loss due to removal of chemisorbed water. Modelling of the mass gain curves using the approaches above could be used in (a) and may be useful in observing transitions in (b).

### 4.2 Carbon Content

It can be observed from that carbon (organic) context was significant in all samples regardless of the context of retrieval, varying from 0.0035 – 0.5198 %wt, Figure 6. As well as this, there does not appear to be notably lower levels present in non-buried samples; this is made clear by the high carbon content in Cau, Por and Bel, and highlight that significant quantities of organic matter should be treated as plausible mass contaminants in all samples used in future RHX dating trials. It is also worth noting that the highest levels were obtained for the samples Por, Etr, Bel, Dow1, Dow2, Rom, Cau (0.52-0.05 %wt) and this will be returned to later. From the TG-MS data, Table 3 and Figures 7-8, it is clear that at heating temperatures from 200-300°C mass loss associated with CO$_2$ removal commences, Event E, corresponding to commencement of the removal of organics (Ford 1967; Rice 1987; Dunham 1992; see Barrett 2015 for a more detailed review of mass loss temperatures associated with firing of ceramics). This removal appears to peak or plateau in the range 400-500°C, Event F, however is not complete until higher temperatures. This is supported by the FTIR data, Figures 15-16 and Table 6, where the presence of organics peaks in region 2800-3000cm$^{-1}$ is still strongly detected even after reheating of powdered samples (<63μm) at 500°C for 18 hours. This is an area
of concern for RHX dating where it is essential that the carbon removed during heating at 500°C equates to that measured as part of carbon content analysis.

From the FTIR identification in the region 3000-2800cm$^{-1}$, four main bands are identified as present in both the non-reheated and reheated samples, Table 6 and Figure 13-15. This region of CH absorption was selected because it is absent of mineral interference and subjectivity issues associated with other regions (see Reeves III 2012). A possible band at 2860-2870cm$^{-1}$, present only in the non-reheated samples, appears to be only an effect of the broad absorption band ranges assigned to samples in Figure 14 because of uncertainty in the peak and is not considered real. The remaining four peaks, summarised below in Table 8, together with infrared bond assignments and organic source interpretations, are made up of the dominant bands, 2845-2860cm$^{-1}$, 2920-2930cm$^{-1}$, and 2950-2975cm$^{-1}$ and a lesser band at 2870-2890cm$^{-1}$. These bands are identified as due to bonds associated with organic matter (CH bonds), specifically aliphatic stretching and bending bonds, Table 8 (identified using Larkin 2011). According to Larkin (2011) the aliphatic stretching bonds often appear as a pair of doublets (in phase, out of phase) and it appears that two sets of doublets are observed in the present work, Table 6, one associated with $R$-$CH_3$ stretching and the other associated with $R$-$CH_2$-$R$ ($R$ is an unspecified group/molecule attached).

Table 8: Table of main FTIR peaks associated with organic matter, together with IR bond assignments, possible organic matter source, and associated references.

<table>
<thead>
<tr>
<th>Wavenumber cm$^{-1}$ (n=non-reheated,r=reheated)</th>
<th>IR bond Assignment (Larkin 2011)</th>
<th>Suspected/Potential Source Organic</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>2. 2880 (n,r) 2870-2890</td>
<td>Aliphatic $R$-$CH_3$ Stretching (in phase)</td>
<td>Not Clear</td>
<td>--</td>
</tr>
<tr>
<td>4. 2960,2970 (n,r) 2950-2975</td>
<td>Aliphatic $R$-$CH_3$ Stretching (out of phase)</td>
<td>Hard Coal, Brown Coal, Coke</td>
<td>van der Marel et al. 1976 Guisnet and Magnoux 2001</td>
</tr>
</tbody>
</table>
Possible organic sources are included in Table 8 together with references for the identification at these bands; however, no source could be assigned to the band at 2870-2890 cm\(^{-1}\) and the sources for the remaining bands are by no means certain. The organic matter in soils is often characterised by a double absorption band at 2855 cm\(^{-1}\) and 2930 cm\(^{-1}\) (e.g. Madari et al. 2006; Vergnoux et al. 2011; Reeves III 2012), yet with no peak at 2950-2975 cm\(^{-1}\) identified by these authors. Fernandes et al. (2010) provide FTIR spectra of the organic matter, humic and fulvic, in some soils and peats; again the double peak at 2855 cm\(^{-1}\) and 2930 cm\(^{-1}\) is evident yet any peak at 2950-2975 cm\(^{-1}\) is absent. Chen et al. (2012) carried out FTIR on a suite of coals varying from peat to anthracite and again the double peaks at 2857 cm\(^{-1}\) and 2927 cm\(^{-1}\) were observed without any third peak apparent (this is supported by similar work by Tian et al. 2010). In this regard, the strong triplet of peaks observed in this study may be more characteristic of a triplet of peaks identified by van der Marel and Beuterspacher (1976) at 2950 cm\(^{-1}\), 2920 cm\(^{-1}\), and 2850 cm\(^{-1}\), found in peat, black peat, brown coal, lignite, hard coal and bitumen. The presence of substances derived from the peat-to-anthracite formation group is certainly possible for raw clays (Dunham 1992), yet it seems unlikely, given the high firing temperatures interpreted for most of the samples (Barrett 2015), that much, if any, of these substances would survive. It is conceivable that, in a reduced (oxygen deprived) firing environment within the kiln (or even within the brick), carbon-rich coke may have been produced and that the major absorption triplet observed in the data could be explained by similar triplet of peaks at 2860 cm\(^{-1}\), 2930 cm\(^{-1}\), and 2970 cm\(^{-1}\), as observed in Guisnet and Magnoux (2001). The use of coke as a fuel in the firing of bricks (in clamps and kilns) is/was not uncommon (Hammond 1981; Brunswick 1990; Pavía and Bolton 2000) and this could also be a source of contamination, particularly post-firing.

From the above, it can not be clarified whether the source of the organics is humic-related substances or derived from part of the peat-to-anthracite formation pathway. From TG-MS, the organics are predominantly being burnt off at temperatures well below the original firing temperature, i.e. less than 500°C and certainly not above 800°C, suggesting that the source of organics removed during the RHX methodology are more likely to be post-firing contamination from burial or air-borne/rain precipitated processes, presumably humic-related substances. That the source of organic matter is likely to be post-firing contamination may be supported by findings from BET analysis, presented in Barrett (2015), where the samples with highest surface area (Table S.3) and pore volume (with the exception of Lan where the carbon content is very low) are also those with the highest carbon contents above (and calcite content, dealt with in Barrett 2015). In Barrett (2015) the BET surface area is also highly correlated \((R^2 = 0.90)\) with the percentage of water removed in the TG-MS heating regime 50-130°C (Figure 5.34 of Barrett 2015), with the greatest quantities of
water loss also occurring for the samples with highest surface area and pore volume, (Figure 5.29 of Barrett 2015). This suggests that high pore volume and surface area may be associated with, and permit, greater uptake of environmental moisture (groundwater, rain), and with it sources of carbon and calcite contamination, supporting an argument that much of the carbon content is derived from post-firing organics of humic origin.

It is not ideal that for each sample only a single estimate of the carbon content was carried out using two measurements of the carbon content of two subsamples, one of which had been previously heated to 500°C. These subsamples were taken from a well-mixed powder of the remaining granules from which the dating subsamples were taken. Hence, the samples used in carbon content analysis are expected to be well homogenised and representative of the dating samples in their carbon content. Nonetheless, variation is to be expected, and with only a single estimate carried out per sample the repeatability of carbon content measurements and the associated uncertainty can not be estimated. As well as this there will be sources of uncertainty in the procedure used for its estimation. However, the author worked under the assumption that the uncertainties in the carbon content, associated with subsample variation and the experimental method, would be minor when compared with that associated with the organic matter to organic carbon ratio; the range of the OM/OC uncertainty used was 1.4-2.5, which equates to a percentage uncertainty of 28% in the mass of organic matter (based on OM/OC ratio of 1.95 used in age estimates), and was considered broad enough to encompass and take account of any sources of uncertainty in the carbon content. These values for the organic matter to organic carbon ratios are based on a survey of research focussed on its estimation for organic matter from a range of environments (aerosol-borne, soil, peats, clays), presented in Table 9. This survey, while not comprehensive, demonstrates that the OM/OC ratio can vary considerably but for all studies falls within the range 1.0-3.0. Notable attention is given to the work of Pribyl (2010) who in a review of 26 existing publications concluded that “a consideration of the possible variation in organic matter composition predicts a range of factor values between 1.4 and 2.5”. This range is used in the present work as an adequate, if not ideal, reflection of the range of likely ratios for contaminants found in ceramics, aerosol-borne or burial-related.

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1 This was constrained by the large number of samples, 36, which needed to be prepared and analysed. Ideally, the carbon content of the subsamples used in mass gain experiments would be captured during heating at 500°C; if conducted on the three subsamples, this would also provide some useful information on the homogeneity of the samples and uncertainty in the carbon content.

2 The XRD, FTIR, and XRF analysis has demonstrated that the two subsamples used in reheated and non-reheated tests were almost identical both mineralogically and in elemental composition.
The mass of organic matter, $m_{OM}$, is a considerable fraction (5-20% typically for better behaved samples on which dating attempts could be made, see Barrett 2015; 2017a) of the total mass loss, $m_{RHX}$, during reheating, Figure 7, and is subtracted (along with loose water, for example) from this quantity in order to obtain the RHX mass used in age estimations, $m_{RHX}$. Therefore, the range of uncertainty used for the OM/OC ratio has a considerable effect on the estimated ages of the samples, Table 4 and Figure 8, causing an uncertainty in the age range of a minimum of 35yrs (for $Cal$ and $t^{14}$) but more typically on the order of at least 100 years for samples where the age estimates are in closest agreement with the known ages (Barrett 2015; 2017a). For example, with the best sample using the $t^{14}$ model the uncertainty associated with the OM/OC ratio leads to an age range for Joy of 321–489 years. If the mid-point in the
This can be re-written as 405±84 years, or a percentage uncertainty of 20%. For the best samples using 

\[ t^{14C} \text{, the age range in } Ann \text{ is 59-182 years, or 120±60 years, a percentage uncertainty of 50%} \]. The effects are quite considerable, then, but do not explain the issue of large ages in most samples.

It is clear then that the presence of organic matter is a serious issue for RHX dating, as previously stated by Numrich et al. (2015), not alone because of the magnitude in which it can occur, but also because it has been found to be present in significant quantities in all samples (only Lan had very low levels), whether the samples were buried during their lifetimes or just exposed to atmospheric conditions. This ubiquitous presence of organic matter raises further questions (in addition to those previously raised by Le Goff and Gallet 2015a; see also the review in Barrett 2015) surrounding good RHX dates obtained in previous studies, where the presence of organic matter was either not examined or treated in a very limited fashion (Wilson et al. 2009; 2012; discussed further in Barrett 2015). It also may be a contributor to poor dates obtained in other studies where the organic matter was not quantified (for example Burakov and Nachasova 2013; Le Goff and Gallet 2014; 2015a). Therefore, in all future RHX dating trials the presence of organic matter needs to be examined out and, if suitable quantification is not possible, a method of its removal is required. Wet chemistry methods may be an option. Numrich et al. (2015) consider the removal of contamination as mandatory and tested several wet chemistry pre-treatments (acid-base leaching, wet oxidation) for the removal of organic carbon. All approaches reduced the carbon content (oxidation, H\(_2\)O\(_2\), proving most successful) but none were successful in providing a complete removal; for this reason the authors recommended pre-treatment cleaning of RHX samples before mass gain tests, with samples still displaying large carbon content following treatment to be discarded. They also stress that these treatments could potentially remove some hydroxyls, or lead to additional hydroxylation. Further work examining the composition (e.g. mineralogy, surface area) and mass loss change in contaminant-free samples with different pre-treatments would be very useful; note that consideration must also be given to the potential effects heating conducted during pre-treatment (Numrich et al. used 60°C for durations of 60-755 minutes) might have on accelerating the rehydroxylation in the ceramics (see the discussion on short term elevated temperature event (STETES) in Barrett 2015).

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3 The age estimate based on an OM/OC of 1.95 was 399yrs old. However, the effects of uncertainties on the age are not symmetric, because the age is proportional to mass\(^4\).
4.3 Mineral Alteration

The differences between the reheated and non-reheated XRD and FTIR spectra showed no appreciable differences (see for example Figure 9) associated with mineral alteration during reheating at 500°C, samples Mac and Bel aside. In associated work (Barrett 2015), it is demonstrated (using XRD, FTIR, petrography) that all samples, with the exception of Etr and Rom, were likely fired at temperature exceeding 850°C, indicated by the presence of high-temperature mineral phases such as cristobalite, spinel and gehlenite. For Etr and Rom there is some evidence (calcite breakdown) suggesting original firing temperatures exceeding 650°C. Therefore, the results generally apply to well-fired samples but are also the case for moderately fired samples. Given the reasonably large set of compositionally diverse samples examined (see Barrett 2015; 2017a; 2017b for details on composition), mass loss associated with mineralogical changes during the reheating stage of RHX is not found to be problematic, aside from gypsum related problems, supporting the findings in Gallet and Le Goff (2015, Figure 1). Also of note is that no mineralogical changes associated with goethite dehydroxylation were observed and, hence, no related mass loss issues (Burakov and Nachosova 2013).

For Mac and Bel, mineralogical changes, Figures 10-11, are associated with the presence and dehydration of hydrated calcium sulfates (gypsum CaSO\(_2\)H\(_2\)O, bassanite CaSO\(_{0.5}\)H\(_2\)O) to anhydrite (CaSO\(_4\)), that occur typically in the temperature range 120-160°C (Dunham 1992). For Mac and Bel, the presence of gypsum was also detected in associated work, precipitated in the pores using petrography and also with sulphur detected using p-XRF ((Barrett 2015). The emission of SO\(_2\) at temperatures exceeding 650-700°C, attributable to gypsum decomposition, is also observed in the TG-MS data, Table 3 and Figure 11.

The presence of gypsum and its dehydration during heating at 500°C for samples Mac and Bel is also associated with a large difference between the S1 mass gains, \(m_{S1}\), of the 130C and 500C components (25°C aging), for example Figure 12 and Table 5, a difference not found to be significant in the other samples (Barrett 2015). There are then two possible issues associated with gypsum. Firstly, there is a mass of water loss during dehydration of gypsum when the samples are heated between 130-500°C; this mass loss will be included falsely in the total fractional mass estimate, \(m_{RHXC}\), resulting in an erroneous increase in the age estimate. Secondly, if the level of physisorption is reduced following the conversion of gypsum/bassanite to anhydrite, then the magnitude of this difference will also be reflected.
as an increase in the total fractional mass estimate (the estimation of the total fractional mass requires that the level of physisorption is the same for both the 130°C and 500°C components).

In the present work, there is no estimate available for the mass loss due to dehydration of gypsum but an estimate for the change in physisorption level is available. This is worth commenting on in terms of the magnitude of the effect it might have on the age estimates in Barrett (2015;2017a), particularly considering that Mac has the highest mass discrepancy (0.00513 and 0.00504, fractional mass for $t^{1/4}$ and $t^{1/n}$ models, respectively; the mass discrepancy corresponds to the quantity of excess mass the RHX mass, $m_{RHX}$, used in age estimates needs to be reduced by to find agreement with the known age) of all samples for both models and Bel is the second highest for the $t^{1/n}$ model (0.00079 and 0.00152 for $t^{1/4}$ and $t^{1/n}$ models, respectively). The experimental estimates of $m_{S1}$ are only approximate (Barrett 2015), but the differences between the 130°C (the larger gain) and the 500°C components, $m_{S1.130} - m_{S1.500}$, are as follows: for Mac, 0.00236 ($t^{1/4}$) and 0.00235 ($t^{1/n}$); for Bel, 0.00355 ($t^{1/4}$) and 0.00385 ($t^{1/n}$). The magnitude is clearly of a similar order to that observed for the mass discrepancies of these samples, particularly for the $t^{1/n}$ model, and therefore, gypsum dehydration, even only considering its effects on physisorption, may be a very significant contributor to issues encountered with the estimated ages of these particular samples. More focussed work on the hydration properties of anhydrite and gypsum in fired clays following heating is required. Screening of samples for gypsum, particularly brick as gypsum can be produced by the reaction of sulphuric acid and calcite (in the composition of the brick or through leaching of surrounding mortars) during weathering (Prentice 1990; Pavía and Bolton 2000), is recommended together with increased examination of methods for its removal that do not interfere with the RHX methodology.
5. Conclusion

Research conducted within the framework of a larger study examining the mass gain behaviour of archaeological ceramics (predominantly bricks) and the application of an RHX dating methodology (component based approach) has revealed new, or provided a better understanding of, mass loss issues that occur during drying or reheating.

In particular, drying of ceramics by conventional (oven) methods requires a prolonged and indefinite period (minimum of months), well beyond that carried out in existing work. Modelling of loose water not removed is a possible solution, however this requires high precision recording of the mass loss curves during drying. Alternatively, if the prolonged drying is due to removal of chemisorbed water, as argued for by the present author, then incomplete drying is not problematic and issues of remaining loose water could potentially be resolved using a component based approach.

The organic matter present in samples is a serious concern for RHX dating and its validation. All samples, regardless of their retrieval context, had significant quantities of carbon present, corresponding to 10-50% of the total mass loss during reheating. This has been attributed to organic matter, most likely humic related substances or coke. On account of large uncertainties in the organic matter to organic carbon ratios (OM/OC), the estimation of the mass of organic matter removed during reheating is problematic; on account of this, the age range estimates of samples are badly affected, often varying from 20-50% of the age of well-behaved samples in associated dating trials. In future dating trials, screening and alternative methods for estimating the organic matter mass are required.

Mineralogical changes during reheating were generally negligible and not considered problematic with the exception of samples where gypsum is present. Dehydration of gypsum during heating leads to an unwanted component of mass loss that is difficult to quantify. As well as this physisorption levels (Stage 1) are altered, requiring some alterations of the equations used in age estimations. Screening of samples is recommended in future work.

The above mass loss issues all have the potential to adversely affect any future RHX dating trials and will require focussed treatment in future work.
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