PROCESSING OF CO₂ SAMPLES COLLECTED USING ZEOLITE MOLECULAR SIEVE FOR ¹⁴C ANALYSIS AT THE NERC RADIOCARBON FACILITY (EAST KILBRIDE, UK)

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ABSTRACT. Cartridges containing a zeolite molecular sieve are used for the field collection of CO₂ from a variety of environments (e.g. atmosphere, soil respiration, evasion from surface waters) for accelerator mass spectrometry (AMS) radiocarbon analysis by the NERC Radiocarbon Facility. Previously, sample CO₂ was recovered from the sieves by heating under vacuum prior to cryogenic purification and graphitization. An additional heating and evacuation stage was used to prepare the cartridge for the next sample. We have recently developed an alternative method using heating and purging with nitrogen gas to remove sample CO₂ from the molecular sieve. An infrared gas analyzer (IRGA) is used to continuously monitor CO₂ content in the purge gas. The preparation of the molecular sieve cartridge for the next use is performed simultaneously and the IRGA reading is used to verify that all sample CO₂ has been removed from the sieve, ensuring insignificant carryover of CO₂ into the next sample. We present the results of a suite of standards of distinctly different ¹⁴C and stable-carbon isotope composition, which we used to confirm the reliability of the new procedure. The new method has considerably reduced the time required to process a single sample from at least 2 hr to less than 40 min, and reduced the minimum sample size by 50%.

INTRODUCTION

A zeolite molecular sieve is a crystalline alumino-silicate with selectivity and adsorption properties that make it particularly useful to trap or purify carbon dioxide (Bol and Harkness 1995). It has been utilized in radiocarbon science from at least the 1960s for sampling atmospheric CO₂ (e.g. Godwin and Willis 1964). It is particularly valuable as an alternative to cryogenic methods for purifying CO₂ on vacuum systems (e.g. Bauer et al. 1992), facilitating its use in the recent automation of laboratory procedures in the preparation of ¹⁴C samples (e.g. Wacker et al. 2010).

At the NERC Radiocarbon Facility (East Kilbride, UK), we use cartridges containing zeolite molecular sieves for the field collection of CO₂ from a variety of environments for ¹⁴C analysis by accelerator mass spectrometry (AMS). For example, the molecular sieve cartridges (MSCs) have been used with a pump-based sampling system (Hardie et al. 2005) to collect CO₂ respired in various environments (e.g. arctic tundra, Hartley et al. 2012; temperate peatlands, Hardie et al. 2009; or evaded from surface waters, Billett et al. 2006, or subsurface pipes, Billett et al. 2012). They have also been deployed as passive samplers, for example, to recover CO₂ for ¹⁴C analysis from the atmosphere (Garnett and Hartley 2010), the permanently waterlogged layers of peats (using hydrophobic filters; Garnett and Hardie 2009), or surface emissions of soils (Garnett et al. 2009).

Hardie et al. (2005) describe the methods we previously used for processing molecular sieve samples, involving heating under vacuum to recover sample CO₂ from the sieve prior to cryogenic purification and graphitization (based on earlier methods described by Bol and Harkness 1995). An additional heating and evacuation stage was used to prepare the cartridge for the next sample. We have recently developed a more efficient method to process CO₂ samples collected using a molecular sieve and to prepare the cartridges for next use. Here, we describe the new method to process samples collected using MSCs and demonstrate the reliability of the method with the results of a suite of standards.

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METHODS

Processing of Molecular Sieve Cartridges

Figure 1 illustrates the vacuum rig used to process MSCs. High-purity N₂ gas (150 mL/min) is introduced to the MSC via a coupling (Colder Products Company, USA). The MSC (containing Type 13X zeolite molecular sieve [BDH, UK] and identical to the design used by Hardie et al. 2005) is placed within a tube furnace (Carbolite, UK) that is initially at room temperature, then heated to 500 °C to release the trapped CO₂. Purge gas exiting the MSC enters the vacuum rig where it is first dried using a slush trap (mixture of dry ice and methylated spirits; −78 °C) and then directed to an infrared gas analyzer (IRGA; PPSystems EGM-4, Hitchin, UK) where the CO₂ concentration is measured. Next, the purge gas passes through 2 liquid nitrogen-cooled traps (−196 °C), retaining the CO₂ and directing non-condensable gases to atmosphere (via a flow gauge). Once the majority of CO₂ has been desorbed from the MSC (when the IRGA readout is <10 ppm), collection of CO₂ in the liquid nitrogen traps is stopped by isolating the traps from the MSC. A vacuum pump is then used to remove non-condensibles from the liquid nitrogen traps, and sample CO₂ is dried for a second time by replacing the second liquid nitrogen trap with a slush trap, and the gas transferred to a calibrated volume to quantify the volume of CO₂ recovered. At the same time as the recovered sample CO₂ is being processed, purge gas continues to flow through the MSC via the IRGA to atmosphere, in order to ensure complete removal of CO₂ from the cartridge (indicated when CO₂ concentration of the purge gas reaches 0 ppm).

Figure 1. Design of the vacuum rig used to process molecular sieve samples by purging with nitrogen gas. A molecular sieve cartridge is shown within the tube furnace. An infrared gas analyzer (IRGA) is used to monitor CO₂ concentration in the purge gas. Small arrows indicate the direction of the purge gas. See Methods for further details.
Test of Laboratory Methods

Tests were performed to verify the new procedure, and especially check for sample carryover (memory effects) since the cartridges had previously been shown not to be affected by fractionation effects (Hardie et al. 2005). A suite of standards (Table 1) of distinctly different $^{14}$C and stable-carbon isotope composition were sampled consecutively on 2 MSCs, and processed with the new procedures.

Table 1 Results for standards used to test the new procedures. Two molecular sieve cartridges (MSC; labeled 1 and 2) were used to consecutively sample CO$_2$ derived from known-age or background standards, after initially being used to sample atmospheric CO$_2$ (assumed to be ~104 pMC). Consensus values for standards reported by Gulliksen and Scott (1995)$^a$ and Boaretto et al. (2002)$^b$.

<table>
<thead>
<tr>
<th>MSC–Run</th>
<th>Source of CO$_2$</th>
<th>Volume CO$_2$ recovered (mL)</th>
<th>$^{13}$C$_{VPDB}$ standard CO$_2$ ±0.1‰</th>
<th>$^{13}$C$_{VPDB}$ recovered CO$_2$ ±0.1‰</th>
<th>$^{14}$C standard CO$_2$ (pMC ±1σ)</th>
<th>Lab code (SUERC–)</th>
<th>$^{14}$C recovered CO$_2$ (pMC ±1σ)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1–1</td>
<td>Atmosphere</td>
<td>5.8</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>1–2</td>
<td>Iceland spar calcite</td>
<td>8.9</td>
<td>&gt;98</td>
<td>+1.6</td>
<td>+1.4</td>
<td>Background</td>
<td>32517</td>
</tr>
<tr>
<td>1–3</td>
<td>TIRI Barley mash</td>
<td>2.6</td>
<td>&gt;99</td>
<td>–26.8</td>
<td>–26.6</td>
<td>116.35 ± 0.0084$^a$</td>
<td>32521</td>
</tr>
<tr>
<td>1–4</td>
<td>FIRI Belfast cellulose</td>
<td>2.9</td>
<td>&gt;95</td>
<td>–23.6</td>
<td>–23.9</td>
<td>57.22 ± 0.04$^b$</td>
<td>32523</td>
</tr>
<tr>
<td>2–1</td>
<td>Atmosphere</td>
<td>6.1</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>2–2</td>
<td>TIRI Barley mash</td>
<td>9.0</td>
<td>&gt;98</td>
<td>–26.8</td>
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<td>116.35 ± 0.0084$^a$</td>
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<td>+2.7</td>
<td>+2.2</td>
<td>Background</td>
<td>32522</td>
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<td>&gt;97</td>
<td>–23.6</td>
<td>–23.8</td>
<td>57.22 ± 0.04$^b$</td>
<td>32516</td>
</tr>
</tbody>
</table>

Standard gases had been initially prepared to CO$_2$ using routine laboratory procedures (combustion/hydrolysis) and stored in sealed glass tubes. To collect each of the standard gases on a MSC, the glass tube containing the standard CO$_2$ was placed in a 1-L air-tight chamber (“Kilner” jar; John Lewis, UK). This chamber was connected to the molecular sieve sampling system described by Hardie et al. (2005) using couplings (Colder Products Company, USA). Atmospheric CO$_2$ was first removed from the chamber by scrubbing with soda-lime, and then the standard CO$_2$ released by breaking the glass tube against the side of the chamber. An air pump (Pulse Pump III, MiDan Co., Chino, California, USA; flow rate ~500 cc/min) was used to circulate chamber air in a closed-loop via a MSC, enabling trapping of the standard CO$_2$. The CO$_2$ trapped on the molecular sieve was then recovered using the new procedures, the MSC recharged and reused for collecting the next standard.

Aliquots of the recovered CO$_2$ were analyzed for $\delta^{13}$C and $^{14}$C concentration. $\delta^{13}$C was determined on a dual-inlet isotope ratio mass spectrometer (VG Optima, UK) at the NERC Radiocarbon Facility and expressed in per mil relative to the Vienna-PDB reference standard. An aliquot of the recovered CO$_2$ was converted to graphite using Fe-Zn reduction (Slota et al. 1987) and passed to the Scottish Universities Environmental Research Centre (SUERC) AMS Laboratory for $^{14}$C measurement. $^{14}$C results were normalized to a $\delta^{13}$C of –25‰ and expressed as pMC following convention (Stuiver and Polach 1977).
RESULTS

Measurements of the CO₂ concentration in the purge gas taken by the IRGA show that CO₂ was rapidly released from the molecular sieve when heated (Figure 2). In fact, it was observed that in some cases within the few minutes that it took for the furnace to reach the target temperature of 500 °C, the CO₂ concentration in the purge gas had already started to decline, indicating that the majority of the CO₂ had been released at a slightly lower temperature. CO₂ concentration fell to 0 ppm (and therefore the MSC recharged for next use) in all samples within less than ~40 min from the commencement of heating the sieve.

The volume of CO₂ recovered from the MSCs processed with the new methods was 95–100% of that expected based on the volume of the standard gas used (Table 1). In all cases, the ¹⁴C concentration of the recovered CO₂ was within measurement uncertainty (2σ) of the reference values for known-age standards (Table 1). Background (¹⁴C-dead) standards had ¹⁴C contents (0.34 ± 0.01 and 0.40 ± 0.01 pMC) only slightly higher than the laboratory background for combusted samples, and were considerably lower than background standards processed using the previous molecular sieve procedure (Hardie et al. 2005). All δ¹³C values were within measurement uncertainty (2σ) with the exception of 1 result (sample 2–3, which was just >2σ).

DISCUSSION

Several studies have previously demonstrated the reliability of zeolite molecular sieve for trapping and purifying CO₂ in samples for ¹⁴C analysis. Some of these studies applied heat and vacuum to recover the sample CO₂, although Wacker et al. (2010) showed that purging with helium efficiently removed adsorbed CO₂ (using N₂ as the purge gas could result in the condensation of N₂ in liquid nitrogen traps [Bauer et al. 1992], but this is not something that we have observed using our method). When incorporated within a vacuum rig system, Bauer et al. (1992) successfully demon-
strated the use of a purge gas approach for processing 14C samples with molecular sieve, as an alternative to cryogenic trapping with liquid nitrogen.

The vacuum and heating procedure previously used at the NERC Radiocarbon Facility for processing our field-collected samples had been shown to be reliable for samples of ~10 mL CO₂ through testing with a suite of 14C and stable-carbon isotope standards. We used a similar testing approach, but with smaller volumes (2.6–9.0 mL) to test the new procedures. It should be noted that the cartridges and sampling system used to trap the CO₂ were almost identical to those used by Hardie et al. (2005); therefore, essentially it was only the new procedures that were being tested.

The results in Table 1 show that the new procedures are reliable for the processing of CO₂ samples collected using MSCs and do not result in significant memory effects, contamination, or isotopic fractionation. Despite consecutively processing samples of distinctly different 14C concentration on the same MSC, the 14C results remained within measurement uncertainty (<2σ) of consensus values. The 14C concentration of background standards (0.34 ± 0.01 and 0.40 ± 0.01 pMC) was only slightly higher than the background associated with routine combustion methods at the laboratory. The results for background standards were also considerably lower than when using the previous method (1.23 ± 0.02 and 1.28 ± 0.02 pMC; Hardie et al. 2005) despite the much smaller size of most of the samples processed using the new procedures. The fact that sample recovery for each of the CO₂ standards was almost 100% further demonstrates the reliability of the new procedures, indicating that no significant quantities of CO₂ remained on the molecular sieve following charging.

The results for test standards demonstrate the reliability of the new procedures, and while we consider that there is still potential for further improvement and refinement (e.g. further reduction in background and/or sample size), the new procedures bring significant benefits. Firstly, the time to process a sample using the new procedures is less than ~40 min, compared with the ~2.5 hr typically required to process a single sample using the previous method. Part of the saving in time is a result of being able to complete the recharging of the MSC at the same time as the recovered sample is still being processed (e.g. aliquoted). However, the greatest saving in time is a result of the rapidity that CO₂ is desorbed from the molecular sieve when purged (Figure 2). Secondly, the new procedures enable the minimum sample size to be processed to be considerably reduced; results from our tests indicate that the new method is reliable for samples with as little as 2.6 mL of CO₂ (compared with the recommended 5–10 mL of CO₂ associated with the earlier method). Finally, incorporation of an IRGA into the purge gas stream allows for a much greater control when processing samples, for example, reliably informing when the molecular sieve has been completely desorbed of sample CO₂.

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