CAPABILITIES OF THE NEW SUERC 5MV AMS FACILITY FOR 14C DATING

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ABSTRACT. A new National Electrostatic Corporation (NEC) 5MV accelerator mass spectrometer became operational at the Scottish Universities Environmental Research Centre (SUERC) in July 2002. It has 2 Cs sputter negative ion sources: a 134-sample source (S1) for the routine measurement of all species, and a hybrid source (S2) with 40 spaces for radiocarbon measurements with either graphite or CO_2 samples. A number of performance tests on graphite samples have been carried out on both sources. A precison of better than 0.3% is feasible for modern samples on a routine basis. The ¹⁴C background of the machine and the graphite preparation process blank are 0.04 ± 0.01 and 0.16 ± 0.05 pMC, respectively, indicating that ¹⁴C dating back to ~50 kyr BP is possible. The normalized ¹⁴C values for a series of reference materials agree well with the IAEA, TIRI, and FIRI consensus values. Routine measurement of ¹⁴C has been underway since May 2003. Preliminary results of performance tests on the CO_2 gas ion source are also reported.

INTRODUCTION

A new NEC 5MV accelerator mass spectrometer (AMS) was installed at the Scottish Universities Environmental Research Centre in July 2002 (Freeman et al., forthcoming a, b). The SUERC spectrometer is equipped with 2 ion sources. One source (S1) accommodates up to 134 samples and is intended for the routine measurement of all species (10 Be, 14 C, 26 Al, 36 Cl, and 129 I), while the other 40-sample source (S2) is a hybrid ion source for radiocarbon measurements with either graphite or CO_2 samples. This paper presents the results for performance tests on graphite samples using both ion sources and also includes preliminary results for performance tests on CO_2 samples.

¹⁴C Performance Tests on Solid Samples

A number of performance tests for ¹⁴C have been performed for the SUERC AMS using the 2 ion sources. The tests have included the study of parameters such as precision, accuracy of normalization, background levels, as well as sample lifetime and throughput. The graphite targets were prepared by the 2 SUERC-hosted ¹⁴C laboratories, NERC lab and SUERC lab, based on the method given by Slota et al. (1987). Each target is composed of 1–2 mg C mixed with 1–2 mg Fe powder. The injection energy of the extracted negative ions was 66 KeV. The measurements were performed at a terminal voltage: 4.5 MV; charge state: +4; total ion energy: 22.5 MeV; and Ar stripper gas pressure: 7–8 Torr. Each sample was scheduled for 5 measurements. For an individual measurement, the modern samples (e.g. oxalic acid and barley mash) were determined until statistical uncertainty reached 0.5% (40,000 ¹⁴C counts), while the background graphite samples were counted for 9000 cycles (about 15 min). However, some samples were completed early if an assigned target value, based on the mean standard deviation of the 3 most recent measurements (0.5% for the modern samples and 10% for background samples), was reached.

For data reduction, an off-line data reduction program calculated 4 values from the measurement data: (1) mean sample isotopic ratio (a weighted average); (2) statistical uncertainty; (3) standard deviation of the mean (SDOM) uncertainty of isotopic ratios; and (4) χ^2 statistics. The program uses the χ^2 statistics to determine if the scatter of the measured isotopic ratios about the average is con-

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sistent with the counting statistic uncertainties of the measurements. Briefly, a sample passes the χ^2 test if its χ^2 statistics are less than or equal to the 5% χ^2 value which varies with degree of freedom, and fails the χ^2 test if its χ^2 statistics are greater than the 5% χ^2 value. For samples that pass the χ^2 test, the program sets the sample ratio equal to the weighted average of the measurement ratios and its uncertainty to the statistical uncertainty. However, for samples that fail the χ^2 test, the simple, not the weighted, average of the measurement ratios and SDOM are used for the sample ratio and its uncertainty, respectively, in subsequent calculations. Moreover, in this study, δ^{13} C values and uncertainties calculated from measured 13 C/ 12 C ratios were used for isotope fractionation correction for 14 C activity or age calculations.

Acceptance Tests

Three oxalic acid standards (OxII) and 3 barley mash samples were used for precision acceptance tests on source S2 (Table 1). In this batch, the $^{12}C^{-}$ currents were between 50 and 58 μ A, with an accelerator 4+ particle transmission of 56%. The standard deviation (1 σ) of the $^{13}C/^{12}C$ and $^{14}C/^{12}C$ ratios of the individual OxII sample was 0.2–0.4% and 0.2–0.3%, respectively. The latter is identical to the statistical uncertainty of 0.2–0.3%. Averaging the 3 OxII samples gives a precision of 0.15% and 0.13% for the $^{13}C/^{12}C$ and $^{14}C/^{12}C$ ratios, respectively. These results showed better than 3‰ precision on the individual and average values. In the case of the barley mash samples, precision on the $^{13}C/^{12}C$ and $^{14}C/^{12}C$ ratios of the individual samples was 0.2% and 0.4%, respectively. These 3 samples yielded better than 3‰ precision not only on the average $^{14}C/^{12}C$ ratio (0.26%) but also on the normalization value (0.27%). It should be pointed out that a repeat of the test at high currents (about 80 μ A), but with reduced accelerator 4+ particle transmission (50%), gave similar results.

The same performance tests were also carried out on source S1. The negative currents extracted from S1 source were 30–35 μ A, slightly lower than those from S2. The average accelerator 4+ particle transmission was 56%. Following the same experimental conditions used in S2, the S1 source gave similar precision for the $^{14}\text{C}/^{12}\text{C}$ and $^{13}\text{C}/^{12}\text{C}$ ratios as observed in S2 source. Therefore, the acceptance tests demonstrated that our new AMS system can perform high-precision measurements of $^{14}\text{C}/^{12}\text{C}$ and $^{13}\text{C}/^{12}\text{C}$ ratios.

System Background

A natural graphite sample (Alfa Aesar graphite powder 100 mesh with a purity of 99.9995%) was measured to assess machine 14 C background, while doublespar (TIRI F) and an interglacial wood (BK-ow) were used to assess system contamination. These materials were always placed adjacent to an OxII or modern sample to monitor any possible cross-contamination between the sample targets. The observed 14 C/ 12 C ratios of Alfa Aesar graphite were generally less than 5×10^{-16} , corresponding to 0.03 pMC (Table 1), which is equivalent to a 14 C age of 64 kyr. This result clearly shows that machine background and source cross-contamination are negligible.

A large volume of CO_2 was produced from doublespar and interglacial wood to minimize contamination during CO_2 production. From this volume, sub-samples of CO_2 equivalent to 1–2 mg C were graphitized. The doublespar yielded 0.16 ± 0.06 pMC, while the interglacial wood yielded 0.16 ± 0.05 pMC. This indicates that a slight contamination has occurred during the graphite target preparation process. Based on the calculation given by Donahue et al. (1990), a ^{14}C dating limit of 51 kyr was obtained.

Materials	¹² C-	¹⁴ C counts	¹³ C/ ¹² C (× 10 ⁻²)	δ ¹³ C (‰)	¹⁴ C/ ¹² C ^b (× 10 ⁻¹²)	¹⁴ C (pMC)
Materials	(µA)	(atoms)	(* 10 -)	(700)	(* 10 12)	(pivic)
Oxalic acid	50	121,338	1.0641 ± 0.0021		1.2094 ± 0.0035	
Oxalic acid	52	121,941	1.0667 ± 0.0045		1.2074 ± 0.0035	
Oxalic acid	53	248,613	1.0643 ± 0.0021		1.2063 ± 0.0025	
Mean for above 3 samples			1.0650 ± 0.0016	-17.8 ± 1.5	1.2077 ± 0.0016	134.07 ± 0.27
Relative deviation (%)			0.15		0.13	
Statistical uncertainty (%)		0.14				
Barley mash (BBM-48)	50	122,202	1.0557 ± 0.0019	-26.4 ± 1.4	1.0426 ± 0.0043	115.78 ± 0.60
Barley mash (BBM-49)	55	122,557	1.0587 ± 0.0023	-23.5 ± 1.1	1.0474 ± 0.0039	116.31 ± 0.56
Barley mash (BBM-50)	58	122,717	1.0584 ± 0.0021	-23.8 ± 1.5	1.0426 ± 0.0044	115.77 ± 0.61
Mean for above 3 samples			1.0573 ± 0.0014	-24.6 ± 1.6	1.0442 ± 0.0028	115.95 ± 0.31
Relative deviation (%)			0.13		0.26	0.27
Alfa Aesar graphite	50	134	1.0781 ± 0.0016		0.00030 ± 0.00003	0.034 ± 0.003

Table 1 Precision of ¹⁴C acceptance test measurement on 19 May 2003 (Source S2)^a.

¹⁴C Accuracy

A series of well-characterized samples were measured to assess the accuracy of the 14 C measurements. These included IAEA C6 (ANU sucrose) and barley mash (TIRI A) to assess accuracy and precision of modern samples, while IAEA C2, C5, C7, C8, Belfast cellulose (FIRI I), and 96 humin (in-house standard) provided samples with a range of ages for similar tests. The results showed that most of the individual measurements agreed with the consensus values within the 1- σ uncertainty margin. In Table 2, average measured 14 C values are compared with the consensus values. It is clear that the relative 14 C age differences between the measured and the consensus values are all in the range from -0.4 to 0.5%.

Table 2 Normalization of reference materials.

			Measured ¹⁴ C		Consensus ¹⁴ C ^a		
Code	Materials	Nr of samples	Activity (pMC)	Age (BP)	Activity (pMC)	Age (BP)	Relative difference (%)
IAEA C6	Sucrose	5	150.31 ± 0.39	_	150.61 ± 0.11	_	-0.20
TIRI A	Barley mash	25	116.35 ± 0.52	_	116.35 ± 0.0084	_	0.00
IAEA C7	Oxalic acid	3	49.65 ± 0.29	_	49.54 ± 0.13	_	0.22
IAEA C2	Travertine	3	41.13 ± 0.27	_	41.14 ± 0.03	_	-0.02
IAEA C5	Wood	4	23.15 ± 0.11	_	23.05 ± 0.02	_	0.43
IAEA C8	Oxalic acid	3	15.11 ± 0.08	_	15.03 ± 0.18	_	0.53
FIRI I	Belfast cellulose	16	_	4494 ± 35	_	4485 ± 6	0.20
96H	Humin	18	_	3365 ± 37	_	3379 ± 66	-0.41
TIRI F	Doublespar	41	0.16 ± 0.06	52,000	0.180 ± 0.006	_	_
BK-ow	Interglacial wood	23	0.16 ± 0.05	52,000	_	_	_
Alfa Aesar	Natural graphite	22	0.04 ± 0.01	63,000	_	_	

^aConsensus values for IAEA C2, C5, and C6 are from Rozanski et al. (1992), IAEA C7 and C8 from Clercq et al. (1998), TIRI A and TIRI F from Gulliksen and Scott (1995), and FIRI I from Scott (2003).

In addition to the performance tests listed above, δ^{13} C values measured with the SUERC AMS were compared with those determined using dual inlet mass spectrometers (VG OPTIMA and MICROMASS SIRA 10). AMS δ^{13} C values were obtained from the AMS-measured 13 C/ 12 C ratio of the sample normalized to the measured 13 C/ 12 C ratio of the OxII standard materials. Figure 1 shows a comparison of δ^{13} C values from the 2 methods for a typical batch of measurements. In this batch, the

^aNo data blocks were rejected.

 $[^]b\text{No}$ background was subtracted and isotope fractionation was corrected using AMS $\delta^{13}\text{C}.$

AMS δ^{13} C values for 102 of the total of 104 samples agreed with the dual inlet MS δ^{13} C values within $\pm 5\%$, which brings about a maximum of ± 40 yr difference when applying the δ^{13} C correction to the ¹⁴C/¹³C ratio in ¹⁴C age calculation. At present, the reason for the difference is not quite clear and more precise measurements over a long term need to be done; however, we expect that the AMS δ^{13} C values can potentially be used for isotopic fractionation correction of the 14 C ages, as has previously only been used in very a few AMS laboratories (e.g. Bonani et al. 1987).

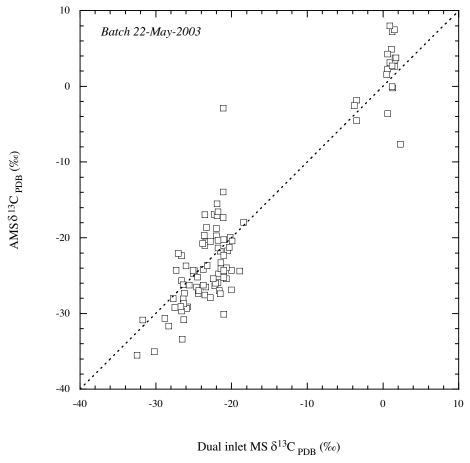


Figure 1 Comparison of δ^{13} C values measured with the SUERC AMS system and dual inlet isotope mass spectrometer (IRMS)

PRELIMINARY PERFORMANCE OF THE GAS ION SOURCE

Although the full performance tests on the gas ion source have not yet been completed, a brief description of the gas ion source with the preliminary performance results are reported as follows. The CO₂ sample is introduced into the 40-sample ion source (S2) by the gas handling system (Figure 2). This handling system uses a manifold and gas flow control system, which applies in 10 individual gas sample containers (bellows). As the bellows (116 cm³) are much larger in volume than the manifold (9 cm³), it can be ensured that little of the gas is wasted when the common part of the manifold is evacuated before another gas sample is introduced. The manifold is connected to the source by a 0.25 mm i.d. capillary of 1.2 m length, giving a constant gas flow for a given pressure. The pressure is measured with a pressure transducer (400 mbar) and this is used to regulate the bellows' volume.

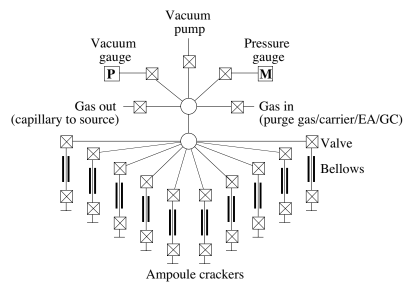


Figure 2 Diagram of the gas handling system

The sample holder used for the gas sample in the ion source is modified from that used for a solid sample by inserting a piece of titanium into the sample holder. The holders are fixed in the sample wheel where the gas tube is pressed on the back of the sample holder. The CO₂ is directly fed through the gas tube to the holder where it is passed over and adsorbed onto the titanium; sputtering by the Cs beam generates negative ions of carbon and oxygen. The targets become sufficiently contaminated with the carbon from the gas so that they must be replaced for each sample.

In operation, gas flow and current stabilization were firstly performed in order to keep the source performance the same for each sample. A typical 10 μA of C^- could be extracted. To determine the CO_2 to C^- efficiency, we used about 250 μg of carbon for a 2-hr measurement, stabilized at 6 μA . This represents an overall efficiency of about 2% for negative ion production. The background beam current, however, was less than 40 nA. These values are comparable to previous reports (e.g. Middleton et al. 1989; Ramsey and Hedges 1997).

In the first measurement with the $^{12}C^-$ currents ranging from 7 to 10 μA and an average accelerator 4+ particle transmission of 57%, the standard deviations (1 σ) of the $^{14}C/^{12}C$ ratios for a single sample, 5 measurements for OxII and doublespar, and 3 for barley mash, were 0.6%, 18%, and 1.1%, respectively, which is comparable to the statistical uncertainties. On the other hand, the precision on the $^{13}C/^{12}C$ ratios for the 3 samples was 0.2–0.3%. Encouragingly, these first results imply that precision for an individual sample is comparable between graphite and the gas source.

CONCLUSION

Since May 2003, the fully-automated and high-throughput SUERC AMS system has been used routinely for ¹⁴C dating. Measurements with modern samples have shown that a precision of better than 0.3% is obtainable. The levels of background for the AMS machine together with the sample preparation process are generally 0.16 pMC, indicating a ¹⁴C dating limit of 51 kyr. Further performance tests on the gas ion source are in progress.

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