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To cite this version:
Yoann Fagault, Thibaut Tuna, Frauke Rostek, Edouard Bard. Radiocarbon dating small carbonate samples with the gas ion source of AixMICADAS. Nuclear Instruments and Methods in Physics Research Section B: Beam Interactions with Materials and Atoms, Elsevier, 2019, 10.1016/j.nimb.2018.11.018. hal-02078529

HAL Id: hal-02078529
https://hal-amu.archives-ouvertes.fr/hal-02078529
Submitted on 25 Mar 2019

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Radiocarbon dating small carbonate samples with the gas ion source of AixMICADAS

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Abstract

Since the beginning of AMS, 14C analyses in foraminifera have been widely used to date deep-sea cores and to estimate reservoir ages of surface and deep ocean layers. Classical AMS requires acid hydrolysis of carbonate samples followed by graphitization of solid targets. For dating very small carbonate samples, an alternative route is to bypass the graphitization step by means of a CO2 gas ion source coupled to a carbonate hydrolysis system.

We report tests and developments performed over 3 years with a carbonate handling system (CHS) coupled to AixMICADAS. Gas source AMS measurements were performed after different pretreatments applied to various carbonate samples including blanks, reference materials, corals and foraminiferal samples ranging from about 5 μg to 100 μg of carbon. In parallel, the CHS coupled to the automated graphitization system (AGES, IonPlus AG) was used to develop a similar leaching procedure for carbonates dated with solid targets. The CHS-AGE3 system can be used efficiently and precisely for samples larger than 300 μgC and less precisely down to 100 μgC by applying a constant contamination correction.

The gas method was thoroughly investigated in order to characterize its performances and limitations. Based on measuring numerous IAEA-C1 blank samples pretreated with hydrochloric acid, an equivalent age of about 52,000 yr BP is routinely achieved, though an even lower background close to 55,000 yr BP is reached when the sample is flushed to suppress the memory effect from sample to sample. The long-term reproducibility is about 7% based on replicated analyses of a modern coral and the accuracy of the gas method was also confirmed by measuring coral and foraminiferal samples previously dated by conventional AMS. The background correction is negligible for samples down to 30 μgC and can be effectively corrected for samples down to 10 μgC by applying a constant contamination correction though the recommended samples size is between 30 μgC and 100 μgC to achieve a reasonable precision.

Our methods were applied to date planktonic foraminifera from two deep-sea cores. Several species with different visual aspects were tested for sample sizes ranging from single shells (ca. 10 μgC) to standard size sample (ca. 100 μgC) for the gas source as well as larger samples (ca. 500 μgC) measured as graphite targets. The dataset does not reveal any large age bias, but illustrates how gas CO2 measurements on small foraminiferal samples could improve the reliability of deep-sea core dating by providing key information on sample heterogeneities due to sediment mixing and diagenesis.

1. Introduction

Radiocarbon analysis of carbonate samples is widely used in paleoceanographic and paleoclimatic studies. For instance, dating planktonic samples is extensively used to establish the chronology of deep-sea sediment cores (e.g. [1]) and 14C age differences between paired benthic-planktonic foraminifera in deep-sea cores can be used to reconstruct surface-deep ocean gradients and provide insight into water mass ventilation (e.g. [2–4]). Reservoir ages can also be reconstructed by comparing planktonic foraminiferal and tephras ages [5–7]. However, this valuable information can only be retrieved from sediment cores where foraminiferal specimens are abundant enough to provide sufficient material (≥1 mg of C) for dating by the conventional method based on the synthesis of graphite targets.
In addition to the sample mass issue, $^{14}$C dating of large samples containing hundreds to thousands of individual specimens can also mask the intra-sample age heterogeneity caused by sediment mixing resulting from physical and biological processes, notably burrowing organisms. This bioturbation has been shown to lead to dating biases when coupled to abundance changes and dissolution of foraminifera [8–13]. Although progress has been made to reduce by an order of magnitude the carbon mass required for graphitization [14–16], it is still a labor-intensive technique, and one which is both prone to contamination effects and which has a limited precision.

An alternative route for dating small carbonate samples is to bypass the graphitization step by using a gas ion source coupled to a hydrolysis unit (e.g. [17–21]). In this paper, we report tests and developments performed over 3 years with the carbonate handling system (CHS, IonPlus AG) coupled to AixMICADAS [22]. More than 1000 AMS measurements were performed with various carbonate samples ranging from ca. 100 μg to 5 μg of C, compared with conventional AMS analyses when enough material was available. This allowed optimizing an analysis protocol including online leaching pretreatment.

2. Method

2.1. Pretreatment and sample hydrolysis

The instrumental set-up for both graphitization and gas measurements have been described elsewhere [19,22]. In short, carbonate samples are weighed and transferred into a 4.5 ml glass vial (Labco Ltd) sealed with a rubber septum and placed in the CHS autosampler. This device is thermostated at 40 °C rather than the 70 °C traditionally used for stable isotope measurements, in order to prevent leaks which can occasionally occur due to the deformation of the polypropylene caps. Each vial is first flushed with helium at 70 ml/min for 3 min by means of a doubled-walled needle, then hydrochloric acid (HCl) is added with concentration and volume adjusted to leach 30% of the initial carbonate mass. Each vial is then gently manually mixed, let to react generally overnight and the CO$_2$ formed during the leaching step is removed by flushing with helium for 3 min at 70 ml/min. Residual samples are then hydrolyzed with 0.06 ml of phosphoric acid (H$_3$PO$_4$ at 85%) and allowed to react for a minimum of 10 min before transferring the first sample of the sequence to the zeolite trap. The released CO$_2$ is pushed with He at 70 ml/min through a water trap (phosphorus pentoxide) into the zeolite trap of the gas source interface system (GIS) or of the AGE3 graphitization system (transfers to the zeolite trap last 2 and 1 min for the AGE and GIS, respectively).

2.2. Graphitization with the automated system AGE3

The pure CO$_2$ is released from the zeolite trap at 420 °C into one of the seven reactors of the AGE3 (IonPlus AG), where it is reduced at 580 °C with graphite and hydrogen (PH$_2$/PCO$_2 = 2.3$) on iron powder (Aifa Aesar, reduced iron 99%, 325 mesh). The reaction time is set to 150 min and the amount of iron is kept constant (3.8–4.2 mg) regardless of the carbon mass. The memory effect from one sample to the next is evaluated at ca. 1% with the AGE3 system connected to the carbonate handling system. In order to minimize this problem, the zeolite trap is first loaded with CO$_2$ of the same age as the sample (artificial mixture prepared from modern and blank CO$_2$ or a split of the CO$_2$ sample to be measured). This so-called PRE-sample is then heated and vented before loading the trap with the real CO$_2$ sample to be graphitized and measured by AMS.

2.3. Measurements with the gas ion source

The pure CO$_2$ is released from the zeolite trap at 450 °C and transferred into the syringe of the GIS where it is mixed with He to obtain a final CO$_2$ concentration of 5% in volume which can be injected into the hybrid ion source of AixMICADAS. The GIS is characterized by a memory effect around 4%, as described in our companion paper (Tuna et al., this issue). The impact of this carry-over can be avoided by loading and purging an aliquot of the CO$_2$ sample to be measured (so-called PRE-sample) and/or by measuring samples in sequence of increasing ages (e.g. following the stratigraphic pile upward).

2.4. AixMicadas parameters and data reporting

The tuning procedure and main operation parameters used in gas and solid configurations are fully described in our companion paper (Tuna et al., this issue [23]). The $^{14}$C data are reported in terms of conventional $^{14}$C age BP and in terms of $^{14}$C/13C isotope ratio after normalization and blank correction ([24–26]). The $^{13}$C/13C ratios are reported using the conventional δ notation by using OxA2 NIST standard.

$^{14}$C blank values are reported without any background or constant contamination while a background correction is applied to standard size samples and an additional correction for the constant contamination is applied to individual foraminifera measurements. Sample masses expressed in this manuscript indicate the residual amount of carbon (μgC) measured from the gas ion source or graphitization system after the pretreatment procedure.

3. Results and discussion

3.1. Pretreatment

Carbonates such as corals and foraminifera are commonly leached with an acid or oxidant prior to $^{14}$C measurements in order to remove secondary calcite or exogenous carbon introduced during sample collection or handling in the laboratory [27–30]. Fig. 1 summarizes the results obtained with different pretreatment methods applied to blank and modern samples.

The relatively large scatter of untreated IAEA-C1 samples compared to leached samples with either HCl or HNO$_3$ suggests that some contamination introduced during sample handling is removed by acid leaching. Although the number of untreated samples is limited, this confirms that acid leaching is useful to get reproducible blanks. The offset for the IAEA-C1 sample treated by the H$_2$O$_2$ oxidative step suggests that a carbonaceous contaminant comes either from the glassware or that it is introduced during the sample handling (e.g. grinding step).

Measurements on foraminifera confirm that untreated samples are less reliable than those leached with either HCl or HNO$_3$. The arithmetic mean of untreated blank foraminiferal samples is $^{14}$C = 0.0027 with a standard deviation (SD) of 0.0003 for n = 3 samples, which is significantly higher than for samples treated either with HNO$_3$ ($^{14}$C = 0.0017, SD = 0.0003, n = 3) or by HCl ($^{14}$C = 0.0018, SD = 0.0004, n = 36). No clear difference is observed for the pretreatments of the standard IAEA-C2 of intermediate $^{14}$C = 0.41. It is probably easy to detect the cleaning effect on blanks. Overall, the results obtained with HNO$_3$ are similar to those with HCl and we have retained this latter acid for our standard leaching procedure. No conclusions can be drawn at present regarding the treatment by the oxidative steps with H$_2$O$_2$ considering the small number of measurements. In addition, this pretreatment method is prone to caps pop-up issues resulting from the pressure build-up caused by the oxygen released by the decomposition of H$_2$O$_2$. 

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3.2. Background contamination for small samples

The quantification of the background contamination for both gas and solid measurements is based on the assumption that samples are mixed with a contaminant of constant mass ($M_c$) and activity ($^{14}C_c$) [e.g. [31-33]]. A modern sample (Porites coral from Moorea) and a blank (IAEA-C1 calcite) of various masses were measured with the hybrid ion source as CO$_2$ gas produced with the CHS and as solid graphite targets prepared with the AGE3. A weighted least squares regression was applied to the results to fit the data (Fig. 2) and determine the constant contaminant parameters ($M_c$ and $^{14}C_c$).

Fig. 1. $^{14}C$ values of gas and solid measurements of standard and blank samples with standard size (residual mass after leaching of $\approx 100 \mu g C$ and $\approx 500-1000 \mu g C$ for gas and solid measurements respectively). (a) Modern coral (Porites from Moorea, Aix-13027), (b) IAEA-C2 reference material, (c) foraminiferal samples picked from a $>100$ kyr sediment layer (marine isotopic stage 5 from a sediment core raised off Morocco) and (d) IAEA-C1 reference material. Data summarized in this figure were measured over a period of two years. The color stand for the different pretreatments applied before the final hydrolysis with H$_2$PO$_4$ (HCl is used routinely, while HNO$_3$ and H$_2$O$_2$ have also been tested). Empty green symbols show individual gas measurements pretreated with HCl, while solid symbols represent the arithmetic mean with the associated standard deviation for both gas (left) and solid (right) measurements. The horizontal light grey bar indicates the arithmetic mean and associated standard deviation of gas samples leached with HCl, while the darker grey bar (panel b) indicates the consensus value and associated error for the reference material IAEA-C2. The results for the modern coral and IAEA-C2 were corrected for a blank value based on IAEA-C1 samples measured in the same magazine. By contrast, the results for IAEA-C1 and blank foraminifera are reported as raw $^{14}C$ without background correction. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Fig. 2. Constant contamination model for (a) gas measurements and (b) solid measurements. Green and blue symbols show $^{14}C$ values with their associated uncertainty (1σ) for the blank sample (IAEA-C1) and modern sample (Porites coral from Moorea), respectively. The red dashed zones indicate the size-ranges of samples that cannot be accurately measured and the red zones designate the size-ranges where samples can be efficiently corrected for the constant contamination. Nevertheless, the precision is markedly impacted due to low ion currents. The white zones represent the recommended carbon mass ranges for optimal precision and accuracy. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
For carbonates with the gas source, the estimated parameters are $M_i = 0.18 \pm 0.04 \mu g$ (2σ) and $F^{14}C = 0.14 \pm 0.04$ (2σ). It should be noted that the data are very well fitted down to a carbon mass of 10$\mu g$ for both modern and blank samples. In addition to the higher influence of the target contamination [34] for samples <10$\mu g$ compared to larger ones, the lower precision and the larger uncertainty for small carbon mass probably explain the scatter for these ultra-small samples. Nonetheless, it is reassuring to see that the background correction is negligible for samples down to 30$\mu g$. This sample mass also allows the full consumption of a Ti target leading to a reasonable precision equivalent to that for standard size ≈100$\mu g$ samples (see our companion paper by Tuna et al. in this issue).

For solid graphite targets, the estimated parameters are $M_i = 1.74 \pm 0.84 \mu g$ (2σ) and $F^{14}C = 0.14 \pm 0.06$ (2σ). This $F^{14}C$ is similar to the CHS-GIS determination and also agree with the recent value determined for the EA-GIS based on the combustion of ultra-small silver cups (see our companion paper by Tuna et al. in this issue).

The $F^{14}C$ of the contaminant with the CHS-GIS is thus lower than the one we found ($F^{14}C = 0.73 \pm 0.11$ and $M_i = 1.45 \pm 0.26 \mu g$) by using the EA-GIS with relatively large silver cups [33] as well as lower than other EA-GIS estimates [31,32]. Higher $F^{14}C$ have also been reported in a recent paper [21] focusing on gas ($F^{14}C = 0.30 \pm 0.04$ and $M_i = 0.68 \pm 0.10 \mu g$) and solid ($F^{14}C = 0.49 \pm 0.07$ and $M_i = 0.94 \pm 0.14 \mu g$) measurements of carbonates with a MICADAS. One possible interpretation is that the contaminating material, assumed as homogenous in the carbon contamination model, is in fact heterogeneous and that the $F^{14}C$ of the residual fraction around 0.2$\mu g$ has a different $F^{14}C$ than the carbon mix forming higher contamination levels around 1$\mu g$. Validation of this hypothesis will necessitate further work.

As seen in Fig. 2, the data are well fitted down to a carbon mass around 100$\mu g$. Although solid samples can be corrected for, it is preferable to graphiteize samples larger than 300$\mu g$ for which the background correction is negligible. In addition, the 12C currents are not optimal and stable for sample masses between 100 and 300$\mu g$ (corresponding to ~20% and ~70% of the typical ion current measured for a 1$\mu g$ sample, respectively). This problem can be reduced by decreasing the amount of Fe catalyst used to produce graphite targets with these small samples.

3.3. Procedural blanks on representative material

Based on 75 measurements of ≈100$\mu g$ aliquots of the reference material IAEA-C1 leached online with HCl, the blank level of the CHS is calculated to be a mean $F^{14}C = 0.0015$, SD = 0.0005 (Fig. 1d and Table 2). This average $F^{14}C$ corresponds to a 14C age of around 52,000 yr BP. This mean $F^{14}C$ value is quite low for a gas source and should be regarded as a conservative estimate because these analyses were mainly performed to purge the CHS before analyzing unknown samples. For instance, an even lower mean value of $F^{14}C = 0.0011$ (≈55,050 yr BP), SD = 0.0003 was obtained from 13 consecutive IAEA-C1 measurements.

Our background is slightly lower than those previously reported [19,21], although a comparison with other MICADAS system is not straightforward due to sample size effects, different pretreatment procedures and improved stripping with He for AixMICADAS compared to the previous system with N2.

As shown in Fig. 1, the procedural blank obtained from gas measurements (≈90$\mu g$) of old (>100 kyr BP) foraminifera leached with HCl is an average $F^{14}C = 0.0018$ (≈50,770 yr BP), SD = 0.0004, n = 36 (average sample mass: 89$\mu g$, SD = 12$\mu g$). The planktonic foraminifera were hand-picked under the binocular microscope from a sediment layer corresponding to marine isotopic stage 5 in a sediment core off Morocco. Although IAEA-C1 and foraminiferal blanks values are both quite low, a small difference of 0.0003 ± 0.0001 can be calculated with foraminiferal blanks consistently higher than IAEA-C1 values.

The same pattern is observed for solid graphite targets of reference materials and foraminiferal samples. Average $F^{14}C = 0.0010$ (≈55,300 yr BP), SD = 0.0002, n = 27 (average sample mass: 900$\mu g$, SD = 190$\mu g$), and 0.0011 (≈54,810 yr BP), SD = 0.0001, n = 2 (average sample mass: 980$\mu g$, SD = 10$\mu g$), were measured on IAEA-C1 and SIRI-K samples (sample K, Sixth radiocarbon laboratory intercomparison, 2013), respectively. These blank levels are lower than those measured on old foraminifera samples from the same core as above with a mean $F^{14}C = 0.0018$ (≈50,980 yr BP), SD = 0.0003, n = 5 (average sample mass: 660$\mu g$, SD = 180$\mu g$). This offset could indicate that a fraction of the surface or intrinsic contamination of these old foraminifera is not completely removed by the leaching procedure.

This agrees with previous observations [35], justifying the measurement of blank levels in samples representative of the same archive (e.g. old foraminifera found in deeper layers of the same deep-sea core). For example, we have re-measured mono-specimen blanks of three foraminifera species from the deeper layers of core MD04-2873 from the Pakistan Margin, which gave a mean $F^{14}C = 0.0018$ (≈50,600 yr BP), SD = 0.0003, n = 3 (Orbulina, G. acutifera and G. ruber, average sample mass: 570$\mu g$, SD = 220$\mu g$) with solid graphite targets and a mean $F^{14}C = 0.0019$ (≈50,170 yr BP), SD = 0.0005, n = 11 (Orbulina and G. ruber, average sample mass: 82$\mu g$, SD = 7$\mu g$) for gas measurements. Both values agree with our previous estimation by conventional AMS [36]. Nevertheless, more work is still needed on sediment cores from different locations and depths in order to characterize the source of the residual contamination in foraminifera.

3.4. Precision and accuracy

The accuracy of the gas method was tested by re-measuring two coral samples previously dated by conventional AMS (Table 1). These well-preserved samples collected offshore Tahiti were used in the frame of the IntCal calibration curve [30]. Both sets of gas replicates measured on ≈100$\mu g$ aliquots are in good agreement for the two corals and the results between the two methods are fully compatible.

More systematic work was performed with gas and solid measurements of a modern coral (MOO, a Porites from Moorea) and the IAEA-

![Table 1](https://example.com/table1.png)

Coral samples from IODP #310 cores offshore Tahiti [28]. Gas measurements are compared with 14C measurements by conventional AMS (French ARTEMIS facility). Samples were corrected with the mean of three IAEA-C1 obtained during the measurement ($F^{14}C = 0.0019$) and a conservative error of ±30% of this value, based on the long-term variability of IAEA-C1 standard size blanks.
C2 standard. The results are summarized and compared in Fig. 1 (panels a & b) and Table 2. The precision for gas and solid measurements can be evaluated from the modern coral. A total of 36 gas samples (over 13 different day-sessions) and 18 solid samples (over 5 different day-sessions) have been measured over a period of two years. Both averages are close to each other, within 2%. As expected, the scatter of the solid measurements (SD = 4%) is smaller than that observed for gas measurements (SD = 7%).

A total of 73 IAEA-C2 gas samples (51 leached with HCl, 2 leached with HNO3, 2 oxidized with H2O2 and 18 untreated) have been measured with the gas ion source over a period of 2 years (over 19 different day-sessions). The data show reproducible results with standard deviations on the order of 10% and 11% of the mean value (leached with HCl and untreated samples, respectively) for this 7 kyr-old sample. This value is in the same order of magnitude than recently reported by the University of Bern (12% and 10% for leached and untreated samples, respectively) [21]. As expected, the scatter for solid measurements is lower than gas measurements, though leached samples seem to be more reproducible (SD = 6% of the mean value, n = 12) than untreated samples (SD = 8% of the mean value, n = 7).

Leached and untreated samples mean values for both solid and gas samples are consistent within one standard deviation with the consensus value (0.414). However, looking at face values, it seems that untreated samples lead to F14C values higher on average than leached samples (6% and 2% for solid and gas, respectively). Although the observed differences are small, they could be linked to a residual contamination of the IAEA-C2 material introduced during the sample handling, as already hypothesized for untreated blanks (see Section 3.1). The fact that similar offsets are observed for solid and gas measurements would exclude a specific effect linked to the gas source.

The mean δ13C value of solid IAEA-C2 samples (−8.0‰, SD = 1.9‰ and −9.1‰, SD = 0.9‰ for leached and untreated samples, respectively) and IAEA-C1 samples (3.5‰, SD = 2.2‰) are compatible within one sigma uncertainty with the consensus value (−8.25 ± 0.31‰ and 2.42 ± 0.33‰ for IAEA-C2 and IAEA-C1, respectively) [35,36]. Gas samples are more reproducible (SD = 1‰) than solid samples but the mean values are systematically shifted toward negative values when compared to solid samples (about −2‰ and −3.5‰ for IAEA-C1 and IAEA-C2 samples, respectively, and −4‰ for the modern coral). Similarly, the δ14C values measured for the different blank samples through the EA + GIS exhibit also a negative shift of around −2‰ when compared to precise and accurate results on solid graphite targets (see our companion paper by Tuna et al. in this issue). By contrast OxA2 samples considered as unknown measured directly from the CO2 standard bottles lead to a mean δ13C of 18.0‰ with an SD of 1.8‰ (n = 132) in agreement with the provided IRMS value (−17.2‰).

The observed δ13C offset is probably due to isotopic fractionation linked to the zeolite trap system of the GIS which are not corrected with our normalization procedure based on the measurement of OxA2 samples from a CO2 bottle. It must be stressed that with the gas ion source, the fate of the CO2 gas is quite different between the OxA2 standard (bottle-CO2 injected directly into the GIS syringe) and the samples (CO2 production with the CHS, drying, adsorption in and desorption from the zeolite trap, mixing of CO2 with He in the GIS syringe).

Normalization with IAEA-C2 carbonate samples has been tested occasionally in order to use the very same procedure for standard and unknown samples. This procedure seems to be efficient to produce less negative δ13C values in better agreement with consensus values for IAEA-C1 and IAEA-C2 samples. However, because of 14C counting statistics, it would be better to normalize the data with a modern carbonate standard rather than with IAEA-C2 (F14C = 0.4). Unfortunately, such a modern carbonate standard is currently not available. This issue was also raised by Gottschalk et al. [21]. In our companion paper by Tuna et al., we propose to use OxA2 samples introduced with the EA in order to normalize data of samples measured through the EA, instead of normalizing with measurements from the OxA2 bottle, which bypass the zeolite trap. Further work will be pursued to develop a similar procedure for carbonates.

In any case, the normalization procedure used so far does not affect 14C/13C ratios as natural and procedural isotopic fractionations are corrected for in the F14C normalization procedure with measured 13C/12C ratios.

3.5. Demonstration of the method on foraminiferal samples

3.5.1. Single foraminifera measurements

In order to test the gas method on small carbonate samples and to study sediment mixing, we have started a measurement program on individual specimens of foraminifera. Fig. 3 shows the individual measurements of 32 single foraminiferal shells (G. truncatulinoides) with masses ranging between 3 and 15 μg C) compared with standard size (±100 μg C) multi-specimen samples of two species (G. truncatulinoides and G. inflata). All samples were sieved and picked under the binocular from the same depth in a sediment core raised off Morocco. A parallel objective of the study was the detection of the contamination by rare modern shells (only two) artificially mixed with the old ones representative of the core depth (see Skinner et al. 2010 [37] also reporting a contamination problem).

Overall, the 32 single foraminifera leached with HCl lead to an average 14C age of 27,790 yr BP with a standard deviation of 1680 yr (represented on Fig. 3 as the green solid square method a). This is larger than the average uncertainty of individual measurements (970 yr). Bioturbation may have contributed to the age heterogeneity. Indeed, this particular deep-sea core has an average sedimentation rate of 14 cm/kyr, which implies that foraminifera over a millennium are mixed in the bioturbation zone (typically on the order of 10 cm).

In addition, the constant contamination model is not ideal for samples smaller than 10 μg C, which also contributed to this large scatter. This non-ideal contamination correction could also be responsible for
Fig. 3. F\(^{14}\)C values corrected for the constant contamination model of 32 single foraminifera samples (the sample represented in red was considered as an outlier) compared to 10 standard size samples from the same depth in deep-sea cores raised off Morocco. Numbers above data points indicate the sample weight in μgC (the mean is 8.3 μg). Letters in red stand for the mean and standard deviation of the 30 single samples leached with HCl (a) corrected for the constant contamination correction, (b) an alternative correction based on two blank foraminifera of similar size (7 and 10 μgC) and, for the sake of comparison, (c) the simple correction based only on standard size blanks. Conservatrive blank errors are propagated to calculate errors bars: ± 40% for blank single foraminifera and ± 25% for standard size samples (based on numerous measurements of MIS-5 foraminiferal samples from this particular core). The horizontal grey bars show the mean and standard deviation for both single and standard size foraminiferal samples pretreated with HCl (for interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

the residual offset observed between the mean result on single foraminifera (27,790 yr BP, SD = 1,680 yr, n = 30) and the mean on standard size (±100 μgC) foraminifera samples (28,150 ± 305 yr BP for G. truncatulinoides and 28,950 yr BP, SD = 380 yr, n = 5, for G. inflata).

An alternative correction with blanks of the same size as the single foraminifera samples, measured during the same sessions, lead to a mean \(^{14}\)C age of 28,220 yr BP, SD = 1710 yr, (Fig. 3, method b) similar to the value calculated with the constant contamination model on single foraminifera and in full agreement with the standard size gas samples. For comparison, the raw data without contamination correction (Fig. 3, method c) give a younger \(^{14}\)C age (27,230 yr BP, SD = 1500 yr).

The gas measurements of standard size samples of foraminifera were also used to test the different chemical pretreatment methods (no treatment, HCl, \(\text{H}_2\text{O}_2\)), but no significant difference was detected with these individual measurements (Fig. 3).

Although this dataset is limited to a specific sediment layer from a single deep-sea core, the data already highlight the advantage of the gas ion source compared to conventional AMS. Indeed, it would have been impossible to study the heterogeneity of this sediment without gas measurement on small CO₂ aliquots (see also [13] for further demonstration). More work is ongoing to compare results of different foraminifera species and size fractions in cores from various locations and depths.

3.5.2. Age models of deep-sea sediments

Constructing an age-depth model requires the dating of numerous discrete core depth intervals. Although valuable age-depth models are routinely based on conventional AMS dating (with ±1 μgC samples) this technique suffers from three main drawbacks. First, it is only applicable when enough foraminifera shells are available, thereby restricting studied locations in the global ocean. Second, many complications such as mixing of the upper sediment layer by burrowing organisms may disturb recorded signals and lead to erroneous interpretation. Third, the diagene-sis is often poorly constrained, which may also lead to incorrect age with large samples [9,38].

As a first test, we analyzed small and large samples from a deep-sea core from the Pakistan Margin in the Arabian Sea (MD04-2876), which is characterized by a high sediment rate (~50 cm/kyr, Bard et al., 2013). The data are shown in Fig. 4, with panels 4a, 4b and 4c showing the details for three particular core depths illustrating how gas measurements on small samples can be used to improve our understanding of sediment mixing and diagenesis.

Several planktonic foraminifera species (G. ruber, G. sacculifer, N. dutertrei, G. bulloides and Orbitolina) were picked under the binocular to provide suitable samples for optimal gas source measurements (±100 μgC representing between 10 and 35 specimens depending on the species). We also distinguished the shells by their visual aspect under the binocular (white, black, opaque, translucent), which may reflect different preservation states or contamination levels by extraneous material. Indeed, the carbonate fraction calculated from the CO₂ pressure in the GIS varies significantly between 25 and 95% of the raw material introduced in the CHS vials. Translucent foraminifera are often considered as particularly well preserved but in this core they may be linked to dissolution as they contain less calcite (69%, SD = 5% n = 4) than opaque or white specimens (80%, SD = 7%, n = 56 and 82%, SD = 7%, n = 4 respectively). Black shells probably contain some pyrite formed in anoxic conditions as the core is located with the Oxygen Minimum Zone. As expected, the calcite content was even lower for these samples (47%, SD = 25%, n = 3).

As can be seen in Fig. 4 (panels a, b and c), the dataset does not reveal any clear age bias between the different species of foraminifera, nor their different aspects. For each of the three depths, the individual gas measurements for one particular species are compatible according to the chi-square test (p = 0.05) and the different species are also in good agreement within one sigma uncertainty of their weighted means. When shell abundance was sufficient, graphite targets were also produced and measured for much larger samples (±350 to 550 μgC). The available solid data (yellow symbols in Fig. 4) confirm the agreement between species and are compatible within one sigma uncertainty with the gas measurements (n = 5). The weighted averages based on the various gas replicates (red symbols in Fig. 4 (panels a, b and c) show that the final weighted error is similar to typical errors for solid measurements (e.g. 25,940 ± 140 yr BP and 25,830 ± 140 yr BP, for gas and solid measurements of Orbitolina from 1450 cm core depth).

The absence of strong discrepancies may seem surprising in the light of the abundant literature on the subject (e.g. [9,13,38]). This may be due to the very high sedimentation rate (50 cm/kyr) and limited bioturbation in these sections of the studied core from the Arabian Sea OMZ. In any case, this preliminary dataset is reassuring concerning
the efficiency of our pretreatment and the reproducibility of the dating protocol.

Fig. 4d provides the age-depth profile for the new $^{14}$C ages measured on planktonic foraminifera with AixMICADAS (both gas and solid measurements). It is compared with the previous age model based on conventional AMS measured at the French ARTEMIS facility [36]. Gas measurements correspond to the weighted mean and associated error from 3 gas measurements. Although more were available for some depths, 3 gas measurements are enough to reach a weighted error comparable to uncertainties of solid graphite analyses.

In general, the new $^{14}$C ages are compatible with the previous ones, except for the 1680 cm core depth for which the new results for gas (29360 ± 340 yr BP) and solid (28880 ± 210 yr BP) are about 1000 yr and 540 yr older than the former age. However, this offset is not much larger than typical error bars in that time range. In addition, the available new measurements were performed on different foraminifera species (G. sacculifer and Orbitolina) than the previous ones (G. ruber). More data will be produced on this particular level to check the reality, size and source of this apparent offset.

4. Conclusion

We pursued technical developments with a carbonate handling system (CHS) coupled to both the gas ion source of AixMICADAS and the automated graphitization system (AGE3) producing solid AMS targets. After testing chemical pretreatments, our preferred method is based on leaching 30% of the sample with HCl, followed by complete hydrolysis of the residual carbonate with phosphoric acid.

For the CHS coupled to the gas source, the average blank on the IAEA-C1 calcite standard corresponds to a $^{14}$C age of 0.0015, SD = 0.0005, n = 75. This mean value is equivalent to a $^{14}$C age of around 52,000 yr BP. An even lower background age of ca. 55,000 yr BP, can be achieved by flushing the CHS to further minimize the sample-to-sample memory.

The procedural blank obtained from gas measurements for foraminifera based on analyzing old (> 100 kyr BP) foraminifera in two deep-sea sediment cores is determined to be an average of $F^{14}$C = 0.0018, SD = 0.0005, n = 47. Foraminiferal blanks are thus slightly, but consistently, higher than IAEA-C1 values. A similar residual difference is observed for solid graphite targets. These observations illustrate the need to measure realistic blanks in order to correct $^{14}$C measurements in a particular sediment core.

The long-term reproducibility of the gas method is about 7% based on replicated analyses of a modern coral and the accuracy was confirmed by dating coral and foraminifera samples previously dated by conventional AMS.

Although the gas source handles samples between ca. 5 µg to 100 µg, it is better to measure samples larger than 30 µg C to optimize precision and reduce the impact of the blank correction uncertainty. Nevertheless, our detailed study of the contamination showed that a correction may be applied for samples down to 10 µg C by assuming a constant contaminant.

Our methods were applied to date planktonic foraminifera from two deep-sea cores. The dataset does not reveal any large age bias between species nor visual aspects of the foraminifera. Measurements on single foraminifera specimens are used to assess the scatter linked to sediment mixing and/or alteration. Although this preliminary study is promising, we plan further tests on foraminifera samples from other sediment cores characterized by different sedimentation rates and dissolution conditions.

Uncited reference

Acknowledgements

AixMICADAS was acquired and is operated in the framework of the EQUIPEX project ASTER-CEREGE with additional matching funds from the Collège de France and the ANR project CARBOTRYDH. We thank Magali Ermini for technical help.

References
