ASSESSING THE BLANK CARBON CONTRIBUTION, ISOTOPE MASS BALANCE, AND KINETIC ISOTOPE FRACTIONATION OF THE RAMPED PYROLYSIS/OXIDATION INSTRUMENT AT NOSAMS

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ABSTRACT. We estimate the blank carbon mass over the course of a typical Ramped PyrOx (RPO) analysis (150–1000°C; $5^{\circ}C \times \min^{-1}$) to be (3.7±0.6) µg C with an Fm value of 0.555±0.042 and a $\delta^{13}C$ value of (-29.0±0.1) % VPDB. Additionally, we provide equations for RPO Fm and $\delta^{13}C$ blank corrections, including associated error propagation. By comparing RPO mass-weighted mean and independently measured bulk $\delta^{13}C$ values for a compilation of environmental samples and standard reference materials (SRMs), we observe a small yet consistent ¹³C depletion within the RPO instrument (mean–bulk: $\mu = -0.8\%$; $\pm 1\sigma = 0.9\%$; n = 66). In contrast, because they are fractionation-corrected by definition, mass-weighted mean Fm values accurately match bulk measurements (mean–bulk: $\mu = 0.005$; $\pm 1\sigma = 0.014$; n = 36). Lastly, we show there exists no significant intra-sample $\delta^{13}C$ variability across carbonate SRM peaks, indicating minimal mass-dependent kinetic isotope fractionation during RPO analysis. These data are best explained by a difference in activation energy between ¹³C- and ¹²C-containing compounds ($^{13-12}\Delta E$) of 0.3–1.8 J × mol⁻¹, indicating that blank and mass-balance corrected RPO $\delta^{13}C$ values accurately retain carbon source isotope signals to within 1–2‰.

KEYWORDS: Ramped PyrOx, blank assessment, kinetic fractionation.

INTRODUCTION

Thermoanalytical instruments such as thermogravimetry (TG) and pyrolysis gas chromatography (pyGC) are frequently used in petroleum geoscience (Peters 1986), biofuels research (White et al. 2011), and soil science (Plante et al. 2009) to monitor the thermal reactivity of organic carbon (OC) contained within environmental samples. Additionally, petroleum geochemists have long coupled thermal analysis methods with isotope ratio measurements in order to investigate the origins and maturity of thermogenic hydrocarbons, leading to the development of techniques such as pyGC-isotope ratio mass spectrometry (IRMS; Galimov 1988; Berner and Faber 1996; Cramer 2004). However, despite their potential to probe the relationship between OC molecular composition, isotope composition, and thermal reactivity, coupled thermalisotope methods have found limited use in other fields of organic geochemistry. Still, preliminary studies analyzing environmental samples indicate that TG coupled with IRMS can yield meaningful trends in stable-carbon isotope (¹²C, ¹³C) composition with temperature (Lopez-Capel et al. 2006; Lopez-Capel et al. 2008). Furthermore, Szidat et al. (2004) and Currie and Kessler (2005) successfully separated and determined the radiocarbon (¹⁴C) content of organic and elemental ("black") carbon fractions in aerosols using a stepped-temperature approach, confirming the possibility that thermal-isotope techniques can be used in tandem with ¹⁴C analysis.

Recently, a novel instrument has been developed at the National Ocean Sciences Accelerator Mass Spectrometry (NOSAMS) facility to determine both the stable and ¹⁴C isotope composition of evolved gases from environmental samples with increasing temperature

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(Rosenheim et al. 2008). This method, termed Ramped PyrOx or RPO, is increasingly being utilized in a host of environments in order to understand the relationship between carbon source, ¹⁴C content, and thermal reactivity (e.g. Rosenheim and Galy 2012; Plante et al. 2013; Rosenheim et al. 2013b; Schreiner et al. 2014; Bianchi et al. 2015). However, a complete understanding of isotope fractionation within the RPO instrument is currently lacking, hindering our ability to accurately interpret evolved-gas ¹³C composition as a carbon source tracer. Additionally, RPO analysis shows promise for improving age-model constraints on carbonate-free sediments (Rosenheim et al. 2013a; Subt et al. 2016), although this application requires that contaminant ("blank") carbon contributions and ¹⁴C mass balance are well constrained. Therefore, the aim of this study is to investigate the blank carbon contribution, isotope mass balance, and kinetic fractionation within the RPO instrument located at NOSAMS.

ANALYTICAL SETUP

The NOSAMS RPO instrumental design is originally described in Rosenheim et al. (2008) and has since been modified to lower contaminant carbon inputs by replacing all plumbing with copper tubing, improve gas flow rates, and improve temperature ramp stability (Plante et al. 2013). In this setup, ultra-high purity (UHP) He gas flows at $32 \text{ mL} \times \text{min}^{-1}$ into a precombusted (850°C, 5 hr) quartz reactor sitting in a two-stage oven containing sample material to be pyrolyzed/oxidized (Figure 1 a,b). He gas is combined with $3 \text{ mL} \times \text{min}^{-1}$ UHP



Figure 1 The NOSAMS RPO instrumental setup: (a.) schematic diagram, (b.) photo of the ovens, and (c.)–(d.) photos of the toggling trap apparatus. Dashed boxes in panel (a.) indicate the regions shown in panels (b.)–(d.).

 O_2 either (1) prior to entering the quartz reactor ("oxidation mode") or (2) downstream of sample material but upstream of a Cu, Pt, and Ni wire catalyst via a reactor side-arm ("pyrolysis mode"). An optimized, combined flow rate of $35 \text{ mL} \times \text{min}^{-1}$ was chosen to minimize transfer time within the system while still allowing sufficient contact time with the wire catalyst and complete cryogenic trapping of CO_2 . During analysis, the lower oven containing the catalyst is held at 800°C to facilitate oxidation of reduced carbon-containing gases to CO_2 , while the upper oven containing the sample is ramped at a user-defined rate with $\approx 5\%$ precision [typically (5 ± 0.2) °C × min⁻¹]. We note that care must be taken when analyzing HCl-fumigated soil/sediment samples (e.g. Plante et al. 2013) as well as marine sediments and dissolved OC, as residual chloride has been observed to interact with and melt the catalysis wire, thus blocking gas flow within the reactor.

After exiting the ovens, water vapor is removed using a dry ice and isopropanol slurry. Gases are then passed into an in-line Sable Systems[®] CA-10 infrared gas analyzer (IRGA) where CO₂ concentration (in parts per million by volume, ppm CO₂) is measured photometrically at 1-s resolution with \approx 5 ppm CO₂ precision in order to generate a plot of temperature versus CO₂ concentration (termed a thermogram). Finally, gases are transferred to a toggling trap apparatus (Figure 1 a,c,d) in which CO₂ is frozen using liquid N₂ while He and O₂ are vented to the atmosphere. At user-defined temperatures, the collecting trap is toggled and CO₂ for each temperature window (termed a fraction) is transferred to a vacuum line, quantified manometrically, and sealed into a precombusted (525°C, 1 hr) 6 mm Pyrex[®] tube containing 100 mg CuO and 10 mg Ag pellets. Following each analysis, tubes are recombusted (525°C, 1 hr) to remove sulfur-containing contaminant gases and CO₂ carbon isotopes are measured following standard NOSAMS procedures (McNichol et al. 1992, 1994a; Pearson et al. 1998). Between each analysis, CO₂ concentration measurements are calibrated using a 2-point calibration curve by plumbing (1) UHP He, and (2) UHP He containing a known CO₂ concentration directly through the IRGA.

RESULTS AND DISCUSSION

NOSAMS RPO Blank Correction

In order to estimate the RPO blank carbon mass and isotope composition, we directly trapped and analyzed CO_2 evolved from empty, precombusted reactor inserts over the typical analytical temperature range (150–1000°C). Although blank carbon contribution is often determined by monitoring deflections from accepted standard reference material (SRM) isotope compositions (i.e. isotope dilution and "modern-dead" methods; Pearson et al. 1998; Santos et al. 2007; Fernandez et al. 2014; Shah Walter et al. 2015), the direct measurement method employed here is better-suited for the RPO instrument for the following reasons:

- 1. Deflections from accepted SRM isotope values are only informative over the narrow temperature range in which the material decomposes, rather than over the course of an entire analysis;
- 2. For stable isotopes, it is possible that kinetic fractionation could overprint isotope deflections due to blank carbon contribution (e.g. Cramer 2004; Dieckmann 2005); and
- 3. Isotope deflection methods are unable to separate blank carbon contributed within the quartz reactor (i.e. time-dependent blank carbon; Fernandez et al. 2014) from that contributed when switching the toggling trap apparatus (i.e. time-independent blank carbon; Fernandez et al. 2014).

reported std. dev. is $\frac{1}{2}$ of the range between values.												
	Mass (µg C)			Flux (ng $C \times {}^{\circ}C^{-1}$)			δ ¹³ C (% vPDB)			Fm		
Toggles	Mean	Std. dev.	n	Mean	Std. dev.	n	Mean	Std. dev.	n	Mean	Std. dev.	n
0	4.0	0.8	4	4.7	0.9	4	-29.0	0.1	1	0.558	0.010	1
2	3.6	0.0	2	4.6	0.0	2				0.595	0.012	1
5	3.4	0.3	2	4.0	0.4	2				0.512	0.013	1
Mean	3.7	0.6	8	4.5	0.7	8	-29.0	0.1	1	0.555	0.042	3

Table 1 NOSAMS RPO blank carbon mass, flux, and isotope composition. For measurements with n = 1, reported std. dev. is instrumental uncertainty. For measurements with n = 2, reported std. dev. is $\frac{1}{2}$ of the range between values.

To address point (3), we calculated the blank carbon mass and ¹⁴C content when the traps were toggled 0, 2, and 5 times at evenly spaced intervals during CO₂ collection between 150 and 1000°C (leading to 1, 3, and 6 collected fractions, respectively). For 2- and 5-toggle experiments, individual fractions were recombined within the vacuum line before transferring to a 6 mm Pyrex tube to keep subsequent steps identical across all experimental conditions. Each experiment was performed in duplicate and the CO₂ mass from each analysis was quantified separately before pairs were combined for ultra-small ¹⁴C analysis (Shah Walter et al. 2015). Results are corrected for the ¹³C/¹²C ratio as measured on the AMS (Santos et al. 2007) and are reported in Fm notation following Stuiver and Polach (1977). We note that Fm reported here is identical to the "¹⁴a_N" notation of Mook and van der Plicht (1999) as well as the "F¹⁴C" notation of Reimer et al. (2004). The 0-toggle experiment was repeated in duplicate for ¹³C analysis using a dual-inlet IRMS as described in McNichol et al. (1994a), and ¹³C/¹²C ratios are reported in δ^{13} C notation (‰ relative to Vienna Pee Dee Belemnite, or VPDB).

Resulting blank carbon mass is independent of the number of toggles throughout the analysis (Table 1), averaging $(3.7 \pm 0.6) \ \mu g \ C \ (n = 8)$ and indicating that the act of toggling the traps contributes a negligible amount of time-independent blank carbon. This is further supported by the near-identical Fm values across experimental conditions (Table 1). We therefore combine measurements from all experiments and calculate an average blank carbon Fm value of $0.555 \pm 0.042 \ (n = 3)$. Because both mass and Fm values are nearly identical across all experiments, we apply the measured 0-toggle blank carbon δ^{13} C value of $(-29.0 \pm 0.1) \ \% o$ VPDB (Table 1) regardless of the number of toggles.

Blank carbon mass calculated here is significantly lower and less variable than that determined for a similar RPO system [c.f. $(12.9 \pm 7.0) \mu g C$; Fernandez et al. 2014], likely due to recent valve and plumbing upgrades on the NOSAMS instrument (Plante et al. 2013). Additionally, photometric measurements suggest that time-dependent blank carbon contribution is not concentrated within any particular temperature range—that is, there exist no distinct peaks within the blank thermograms (Figure 2). Although the mean blank flux appears to drop slightly from $(5.8 \pm 0.7) \text{ ng C} \times °C^{-1}$ when T < 550°C to $(3.1 \pm 1.0) \text{ ng C} \times °C^{-1}$ when T ≥ 550°C, it can nonetheless be reasonably described as constant throughout the analysis within the 95% confidence interval of the manometric measurements (Figure 2).

Dividing the manometric blank carbon mass by the experimental temperature range results in a blank carbon flux of (4.5 ± 0.7) ng C×°C⁻¹ (assuming a 5°C×min⁻¹ ramp rate; Table 1). We therefore correct the mass of carbon in each RPO fraction for blank contribution according to

$$m_s = m_m - \phi_b \Delta T \tag{1}$$



Figure 2 RPO blank carbon flux for a ramp rate of $5^{\circ}C \times \min^{-1}$ as determined photometrically and manometrically. For photometric measurements, absolute CO₂ concentrations were normalized such that the mean value for each analysis is equal to the manometric mean, as small differences in IRGA baseline calibration between analyses leads to large changes in calculated blank flux.

where m_s is the true sample carbon mass, m_m is the measured carbon mass, ϕ_b is the blank carbon flux (in units of mass $\times {}^{\circ}C^{-1}$), and ΔT is the temperature range over which the CO₂ was collected. Here, we proceed using the manometric average ϕ_b value of (4.5 ± 0.7) ng C $\times {}^{\circ}C^{-1}$. However, we note that temperature-specific ϕ_b values listed above could offer slight improvements in blank-corrected mass accuracy, although these results will typically be statistically identical to those using the manometric average value. Additionally, we propagate uncertainty for this correction according to

$$\sigma_{m_s} = \sqrt{\left(\sigma_{m_m}\right)^2 + \left(\sigma_{\phi_b} \Delta T\right)^2} \tag{2}$$

where σ is the standard deviation associated with each subscripted measurement. This assumes that ΔT is known perfectly (i.e. $\sigma_{\Delta T} \equiv 0.0$) and that the uncertainty in m_m and ϕ_b are uncorrelated, which is reasonable given that $m_s \approx m_m \gg \Delta T \phi_b$. Similarly, we treat the measured CO₂ isotope composition as a weighted average of sample carbon and blank carbon, and correct for blank contribution following

$${}^{x}\mathbf{R}_{s} = \frac{m_{m}{}^{x}\mathbf{R}_{m} - \phi_{b}\Delta \mathbf{T}^{x}\mathbf{R}_{b}}{m_{s}}$$
(3)

where ^xR_i is the ^xC/¹²C isotope ratio of component *i* [x = 13, 14; *i* = (s)ample, (m)easured, (b)lank], with ¹³R_i expressed in δ^{13} C notation (% VPDB) and ¹⁴R_i expressed in Fm notation. Lastly, we propagate uncertainty associated with isotope corrections. Because $m_s \approx m_m$, we cancel these where appropriate to avoid large covariance terms, leading to

$$\sigma_{\mathbf{x}_{\mathbf{R}_{s}}} \cong \sqrt{\left(\sigma_{\mathbf{x}_{\mathbf{R}_{m}}}\right)^{2} + \left(\frac{\Delta \mathbf{T}^{\mathbf{x}}\mathbf{R}_{b}}{m_{s}}\sigma_{\phi_{b}}\right)^{2} + \left(\frac{\phi_{b}\Delta \mathbf{T}}{m_{s}}\sigma_{\mathbf{x}_{\mathbf{R}_{b}}}\right)^{2} + \left(\frac{\phi_{b}\Delta \mathbf{T}^{\mathbf{x}}\mathbf{R}_{b}}{m_{s}^{2}}\sigma_{m_{s}}\right)^{2}} \tag{4}$$

For typical RPO fraction CO₂ masses ($\approx 100 \,\mu g$ C) and ΔT ($\approx 100^{\circ}$ C) encountered during sample analyses, blank carbon correction shifts δ^{13} C values by -0.02% (for δ^{13} C = -35% VPDB) to +0.15% (for δ^{13} C = +5% VPDB) and Fm values by -0.002 (for Fm = 0.01) to +0.002 (for Fm = 1.0), within the typical analytical uncertainty of these measurements. While

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¹⁴C content of graphite targets containing as little as $6 \mu g$ C has been accurately analyzed at NOSAMS (Shah Walter et al. 2015), we recommend a minimum RPO fraction mass of $25 \mu g$ C in order to keep blank carbon corrections below 0.5% for δ^{13} C and 0.01 for Fm (assuming $\Delta T = 100$ °C). A spreadsheet for performing all blank correction calculations is included in the supplementary material (Table S1).

Isotope Mass Balance

If sample carbon is completely converted to CO_2 by the end of an analysis and is efficiently transferred to the vacuum line, the mass-weighted mean CO_2 isotope composition of blankcorrected RPO fractions should match independently measured bulk values within analytical uncertainty. To test this, we compare RPO mass-weighted mean compositions with bulk measurements for a range of sample types (SRMs, dissolved organic carbon, fluvial/marine total suspended sediments, soils, and lacustrine/marine sediments). Bulk $\delta^{13}C$ values were obtained either using an elemental analyzer coupled to a continuous-flow IRMS following Whiteside et al. (2011) or on a dual-inlet IRMS after conversion to CO_2 by closed-tube combustion as described in McNichol et al. (1994a). Bulk Fm was measured at NOSAMS following standard preparation methods for each sample type (McNichol et al. 1994b) and uncertainty for each bulk measurement is taken as the measured analytical uncertainty. We calculate RPO mass-weighted mean isotope compositions ($\overline{^{X}s}$) following

$$\overline{^{\mathbf{x}}\mathbf{R}_{s}} = \sum_{j=1}^{n} f_{j}^{\mathbf{x}}\mathbf{R}_{s,j}$$
(5)

where *n* is the total number of CO₂ fractions collected throughout the analysis, f_j is the contribution of fraction *j* to the total mass of CO₂ such that $\Sigma_j f_j \equiv 1.0$, and ${}^{x}R_{s,j}$ is the blank-corrected ${}^{x}C/{}^{12}C$ isotope ratio of fraction *j*. Additionally, assuming that f_j is known perfectly (i.e. since $\Sigma_j f_j$ must equal 1.0 by definition), we estimate the mass-weighted mean isotope uncertainty according to

$$\sigma_{\overline{sR_s}} \cong \sqrt{\sum_{j=1}^n (f_j \sigma_{sR_{sj}})^2}$$
(6)

To test the ability of RPO mass-weighted mean isotope values to predict measured bulk values, we performed orthogonal distance regression (ODR), including uncertainty in both x and y variables, using the SciPy package in Python v3.5. and a weighting factor for each sample that is inversely proportional to the uncertainty in each measurement (Boggs and Rogers 1990; Oliphant 2007). All data presented here are either taken from the literature (Rosenheim and Galy 2012; Rosenheim et al. 2013a) or are originally presented in this study.

Stable Isotope Mass Balance

On average, the RPO mass-weighted mean isotope composition is depleted in ¹³C by $(0.8 \pm 0.9) \%$ relative to bulk measurements (n = 66) independent of RPO analytical conditions (Figure 3), as has been described previously (Rosenheim and Galy 2012; Rosenheim et al. 2013a). To test if residual ¹³C-enriched carbon remaining after RPO analysis could cause this depletion, Rosenheim and Galy (2012) requantified the carbon content of total suspended sediment samples after ramping to 1000°C and determined that only $\approx 0.003\%$ of initial carbon remained. Therefore, for the samples tested therein, Rosenheim and Galy (2012) concluded that low yield could not explain the observed bias. We tested additional potential sources



Figure 3 (a.) Cross-plot of RPO mass-weighted mean versus independently measured bulk δ^{13} C values for all samples in this study in which δ^{13} C data exist and (b.) the same data presented as a histogram of deviations from bulk values ($\Delta \delta^{13}$ C = δ^{13} C_{mean} $-\delta^{13}$ C_{bulk}). Sample abbreviations are as follows: DOC, dissolved organic carbon; TSS, total suspended sediments; SRM, standard reference material.

of this depletion by performing a series of experiments using a CO₂:He calibration gas mixture with known isotope composition [465.5 ppm CO₂ in He, $\delta^{13}C = (-14.9 \pm 0.04)$ ‰ VPDB] as follows:

- 1. Plumbing calibration gas directly into the toggling traps (bypassing the ovens of the RPO system) over a range of flow rates: 15, 35, and 50 mL × min⁻¹;
- 2. Freezing CO₂ from the calibration gas for a range of integration times for each of the flow rates in experiment (1): 1, 5, and 10 min; and
- 3. Plumbing calibration gas through an empty, precombusted reactor insert and collecting CO_2 between 150 and 1000°C, toggling every 170°C for a total of 5 fractions (flow rate = $35 \text{ mL} \times \min^{-1}$, ramp rate = $5^{\circ}C \times \min^{-1}$).

The results of experiments (1) and (2) reveal that, for all flow rates and integration times, the collected CO₂ δ^{13} C value [(-15.0 ± 0.1) ‰ VPDB, n = 9] is statistically identical to the accepted value, indicating that dynamic cryogenic trapping within the toggling traps imparts no isotope fractionation. Furthermore, oven temperature does not appear to affect ¹³C composition, as δ^{13} C values from all fractions in experiment (3) are statistically identical with a mean value of (-15.2 ± 0.04) ‰ VPDB (n = 5). Although this is 0.3‰ depleted relative to the accepted value, this bias is smaller than that observed in most samples within our sample set (i.e. up to 3‰, Figure 3b), suggesting that any fractionation imparted during transport through the hot oven alone cannot cause observed ¹³C depletion.

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However, we note that the mass-weighted mean versus bulk δ^{13} C difference is more pronounced in decarbonated samples containing exclusively OC (mean-bulk: $\mu = -1.0\%$; $\pm 1\sigma = 0.9\%$; n = 60) as compared either to samples containing mixtures of carbonate and OC or pure carbonate SRMs (mean-bulk: $\mu = -0.1\%$; $\pm 1\sigma = 0.5\%$; n = 6). We therefore hypothesize that isotope fractionation during OC degradation within the RPO oven could cause ¹³C depletion, potentially due to incomplete oxidation to CO₂ while reduced carbon-containing gases are in contact with the catalyst wire (Figure 1a). This mechanism is consistent with the results of experiment (3) indicating a lack of temperature dependence on isotope fractionation. We therefore recommend that δ^{13} C values of each RPO fraction *j* within a particular sample can be fractionation-corrected according to the difference between mass-weighted mean and bulk measurements of that sample

$$\delta^{13} \mathbf{C}_{s,j,\text{corrected}} = \delta^{13} \mathbf{C}_{s,j} + \left(\delta^{13} \mathbf{C}_{\text{bulk}} - \overline{\delta^{13} \mathbf{C}_s}\right)$$
(7)

Furthermore, assuming that the covariance between $\delta^{13}C_{s,j}$ for each fraction *j* and the massweighted mean value $\left(\delta^{13}C_{s}\right)$ is small compared to all other variance terms, we propagate uncertainty associated with fractionation correction according to

$$\sigma_{\delta^{13}C_{s,j,\text{corrected}}} \cong \sqrt{\sigma_{\delta^{13}C_{s,j}}^2 + \sigma_{\delta^{13}C_{\text{bulk}}}^2 + \sigma_{\overline{\delta^{13}C_s}}^2} \tag{8}$$

¹⁴C Mass Balance

In contrast to δ^{13} C, mass-weighted mean Fm values typically agree with bulk Fm values within analytical uncertainty across all sample types and analytical conditions (mean-bulk: $\mu = 0.005$; $\pm 1\sigma = 0.014$; n = 36; Figure 4). This can be easily explained because Fm is by definition corrected for the 13 C/ 12 C ratio as measured on the AMS (Stuiver and Polach 1977; Santos et al. 2007) such that any mass-dependent fractionation occurring in the RPO instrument is accounted for. It is additionally useful to compare relative deviations between bulk and RPO mean values, as 14 C content of samples is highly variable. For the samples analyzed here, this equates to an average mean – bulk relative difference of 1.0% with a standard deviation of 3.3% (n = 36), independent of absolute 14 C content of the sample (Figure 4b). This agreement between the mass-weighted mean Fm and bulk Fm values further precludes the possibility that a significant amount of isotopically unique carbon remains unreacted after ramping to 1000°C, and is strong evidence that 14 C mass balance during RPO analysis is robust over the entire range of Fm values found in nature.

Kinetic Fractionation

Finally, we evaluate the kinetic isotope effect (KIE) due to mass-dependent differences in pyrolysis/oxidation rates between each isotope during temperature ramping. If the amplitude of the KIE is significant relative to natural compositional differences, then changes in δ^{13} C values between RPO fractions within a single sample can reflect instrumental fractionation rather than differences in carbon source isotope composition. Quantifying fractionation due to the KIE is therefore critical in order to interpret ¹³C composition as a carbon source tracer. To do so, we measured δ^{13} C values of evolved CO₂ from two carbonate SRMs in high-resolution fashion by toggling every $\approx 20^{\circ}$ C: (1) travertine calcite (IAEA C2; Rozanski et al. 1992), and (2) Icelandic spar [in-house standard; long-term average δ^{13} C = (3.00 ± 0.03) % VPDB]. Because carbonates are chemically and isotopically homogenous, any resulting δ^{13} C variability should follow a predictable, Rayleigh-like fractionation line that depends only on the difference in activation energy (*E*) between the decomposition of ¹³C- and ¹²C-containing molecules (¹³⁻¹² $\Delta E = {}^{13}E - {}^{12}E$; Kwart 1982).



Figure 4 (a.) Cross-plot of RPO mass-weighted mean versus independently measured bulk Fm values for all samples in this study in which Fm data exist and (b.) the same data presented as a histogram of relative deviations from bulk values, in percent $\left[\Delta Fm\left(\%\right) = \frac{Fm_{mean} - Fm_{bulk}}{Fm_{bulk}} \times 100\%\right]$. Sample abbreviations are as follows: DOC, dissolved organic carbon; TSS, total suspended sediments.

We describe the carbonate decomposition rate constant at any temperature [k(T)] by an Arrhenius equation (here written for ¹²C)

$$^{12}k(T) = {}^{12}k_0 \exp\left(-\frac{{}^{12}E}{RT}\right)$$
 (9)

where ${}^{12}k_0$ is the Arrhenius pre-exponential factor for ${}^{12}C$ and *R* is the ideal gas constant. Following Kwart (1982), the KIE at any temperature [KIE(T)] is defined as the ratio of ${}^{12}C$ and ${}^{13}C$ rate constants at that temperature

$$\text{KIE}(\text{T}) = \frac{{}^{12}k(\text{T})}{{}^{13}k(\text{T})} = {\binom{{}^{12}k_0}{{}^{13}k_0}} \exp\left(\frac{{}^{13-12}\Delta E}{R\text{T}}\right)$$
(10)

Equation 10 fundamentally states that, for a given ${}^{13-12}\Delta E$, ${}^{12}k_0$, and ${}^{13}k_0$, KIE(T) decreases with increasing T, indicating that kinetic fractionation within the RPO instrument will be largest for lower temperature components. Furthermore, we can reasonably assume that entropic differences between 13 C- and 12 C-containing molecules are negligible within the carbonate crystal lattice (c.f. Tang et al. 2000). This assumption implies that ${}^{12}k(T) = {}^{13}k(T)$ as T approaches infinity and requires that ${}^{12}k_0 = {}^{13}k_0 = k_0$ (Cramer 2004). Additionally, for each temperature we compute the 13 C composition of the remaining carbonate that has not yet decomposed [13 R_{carb}(T)] as

$${}^{13}\mathbf{R}_{carb}(\mathbf{T}) = \overline{{}^{13}\mathbf{R}_s} \exp\left(\frac{{}^{12}\mathbf{I}(\mathbf{T}) - {}^{13}\mathbf{I}(\mathbf{T})}{\beta}\right)$$
(11)

-	•	-	-	-	
Sample	Analysis type	$k_0 ({ m s}^{-1})$	^{12}E (kJ × mol ⁻¹)	$^{13-12}\Delta E$ (J × mol ⁻¹)	Reference
Travertine (IAEA C2) Icelandic spar	RPO (oxidation) RPO (oxidation)	1.0E + 15 1.0E + 15	326 324	1.8	This study This study
Tarim Basin kerogen Tarim Basin crude oil	Sealed pyrolysis Sealed pyrolysis		218 230	2–234 –52–314	Tian et al. (2007) Tian et al. (2007)
Westphalian coal Individual hydrocarbons	pyGC-IRMS Pyrolysis <i>ab initio</i> modeling	2.4E + 14	230–310 167–500	30–110 15–242	Cramer (2004) Tang et al. (2000)

Table 2 Comparison of k_0 , ¹²E, and ^{13–12} ΔE values for carbonate SRMs in this study with those calculated using various thermoanalytical techniques on petroleum products.

where β is the oven ramp rate, $\overline{{}^{13}R_s}$ is the mass-weighted mean ${}^{13}C$ content of the sample calculated by Equation 5, and ${}^{12}I(T)$ and ${}^{13}I(T)$ are the temperature integrals for ${}^{12}C$ - and ${}^{13}C$ -containing molecules according to Braun and Burnham (1987) (here written for ${}^{12}C$)

$$^{12}I(T) \simeq \frac{RT^2}{^{12}E} {}^{12}k(T) = \frac{k_0 RT^2}{^{12}E} \exp\left(-\frac{^{12}E}{RT}\right)$$
 (12)

Finally, following Cramer (2004), we calculate the predicted ¹³C composition of instantaneously evolved CO₂ at any temperature [$^{13}R_{CO_2}(T)$]

$${}^{13}R_{CO_2}(T) = \frac{{}^{13}R_{carb}(T)}{KIE(T)} = {}^{13}R_{carb}(T)exp\left(-\frac{{}^{13-12}\Delta E}{RT}\right)$$
(13)

Calculating ${}^{13}R_{CO_2}(T)$ requires two inputs in addition to ${}^{13-12}\Delta E$: k_0 and ${}^{12}E$. Here we prescribe k_0 a priori and estimate ${}^{12}E$ for each SRM by minimizing the root mean squared error (RMSE) between predicted first-order decay rates and observed thermograms using a Nelder-Mead algorithm in the SciPy package for Python v3.5. (Table 2; Nelder and Mead 1965; Oliphant 2007). We note that ${}^{13}R_{CO_2}(T)$ is insensitive to our choice of k_0 (Dieckmann 2005; White et al. 2011). For example, assuming a large ${}^{13-12}\Delta E$ value of 100 J × mol⁻¹ for a peak at 700°C, changing k_0 from 10^{10} s⁻¹ to 10^{20} s⁻¹ increases δ^{13} C of the first 1% of evolved CO₂ by only 1%₀ and the first 50% of evolved CO₂ by only 0.2‰. We therefore reasonably choose $k_0 = 10^{15}$ s⁻¹ based on a compilation of literature values [see White et al. (2011) for review]. We then calculate ${}^{13-12}\Delta E$ that best predicts the 13 C composition of all CO₂ fractions for each SRM by minimizing the measured versus predicted RMSE (Nelder and Mead 1965; Oliphant 2007). To accurately compare instantaneous 13 C content predicted by Equation 13 to measured RPO fractions (which integrate over time), we use the CO₂-mass-weighted average temperature for each fraction.

Measured ¹³C composition for both SRMs is consistent with a ${}^{13-12}\Delta E$ value between 0.3 and 1.8 J × mol⁻¹ (Table 2; Figure 5), significantly smaller than literature values for petroleum products using various non-isothermal pyrolysis instruments (Table 2). Therefore, for the SRMs analyzed here, predicted CO₂ δ^{13} C increases by <1% until >>99% of initial carbon has been decomposed (Figure 5). However, we note that, on one hand, calculated ${}^{13-12}\Delta E$ using carbonate SRMs is likely a minimum estimate for environmental samples, as this carbon is already present in a + IV oxidation state, while oxidation of OC could increase ${}^{13-12}\Delta E$. On the other hand, it has been shown that samples with high molecular diversity—as is expected in environmental OC mixtures—exhibit less *apparent* kinetic isotope fractionation than do single



Figure 5 RPO fraction δ^{13} C values for two carbonate SRMs [(a.) travertine and (b.) Icelandic spar] plotted with the predicted δ^{13} C value at each temperature using best-fit $^{13-12}\Delta E$ values from Equation 13 (solid black line). For reference, predicted δ^{13} C values for various $^{13-12}\Delta E$ values are plotted as dashed and dotted lines, while shaded gray regions represent normalized thermograms (unitless). Each RPO fraction is plotted at its CO₂-mass-weighted mean temperature.

compounds such as the carbonates analyzed here (Cramer 2004). Overall, we recommend that a $^{13-12}\Delta E$ range of 0.3–1.8 J × mol⁻¹ is valid for any component within an RPO analysis, and we consequently predict that kinetic isotope fractionation cannot exceed 1.8% during pyrolysis/ oxidation of the first 99% of any sample eluting between 150 and 1000°C. In reality, 13 C enrichment at >>99% combustion will never be observed during RPO analysis, as each fraction typically contains 10–20% of total carbon. We therefore conclude that δ^{13} C variability greater than 1–2% between RPO fractions must reflect differences in source carbon isotope composition.

Furthermore, if kinetic fractionation were driving observed ¹³C variability, δ^{13} C values of evolved CO₂ from all samples should increase monotonically with temperature along a trend that depends only on ^{13–12} ΔE , which is clearly not observed. Rather, the δ^{13} C spread (i.e. max-min) across RPO fractions is highly variable between samples, reaching values as high as 28.8% in carbonate-containing lacustrine sediments and as low as 0.3%



Figure 6 RPO fraction δ^{13} C values for two environmental samples: (a.) decarbonated Ganges River TSS (Galy et al. 2008) and (b.) Hawaiian soil (Chadwick et al. 2007). δ^{13} C values do not show a monotonic increase with temperature, precluding the possibility that δ^{13} C variability in these samples reflects kinetic fractionation. For reference, shaded gray regions represent normalized thermograms (unitless). Each RPO fraction is plotted at its CO₂-mass-weighted mean temperature.

in decarbonated soils. For three carbonate-containing sediments analyzed here, we additionally measured the δ^{13} C value of total inorganic carbon following standard methods (McNichol et al. 1994b) to compare with blank and mass-balance corrected RPO results. For all samples, high-temperature RPO δ^{13} C values agree with those of total inorganic carbon within 1%, further indicating that RPO δ^{13} C values accurately reflect source carbon composition.

Lastly, decreasing δ^{13} C values have been observed with increasing temperature in select samples such as decarbonated Ganges River total suspended sediments and Hawaiian soils (Figure 6), opposite of trends that would depict kinetic fractionation. Rather, this agrees with the interpretation that labile C₃ OC in these environments is replaced by ¹³C-enriched, C₄-derived material (Chadwick et al. 2007; Galy et al. 2008), and is further evidence that measured δ^{13} C

trends reflect differences in carbon source isotope composition. Combined, the RPO δ^{13} C trends from environmental samples analyzed here agree with SRM-based fractionation predictions indicating that kinetic fractionation is small (i.e. less than 1–2‰) in the RPO instrument at NOSAMS.

CONCLUSION

We describe the blank carbon composition, isotope mass balance, and kinetic isotope fractionation within the NOSAMS RPO instrument. Blank carbon mass is significantly smaller than that reported on a similar system (Fernandez et al. 2014) and can be described as a constant flux of (4.5 ± 0.7) ng C × °C⁻¹ (for a 5°C × min⁻¹ ramp rate) with an Fm value of 0.555 ± 0.042 and a δ^{13} C value of (-29.0 ± 0.1) ‰. We find no evidence for significant time-independent blank contribution, likely due to recent valve and plumbing upgrades within the instrument (Plante et al. 2013).

Isotope mass balance on a suite of environmental samples indicates that independently measured bulk Fm is accurately reconstructed using the RPO fraction mass-weighted mean. In contrast, RPO-predicted weighted-average δ^{13} C values are slightly depleted relative to measured bulk δ^{13} C values, especially for decarbonated samples containing exclusively OC. We eliminate the possibility that this depletion is due to low carbon yield or fractionation within the toggling traps. Rather, we hypothesize that this is caused by incomplete oxidation of reduced gases to CO₂ within the oxidation oven and suggest that δ^{13} C of each RPO fraction for a given sample can be mass-balance corrected using the difference between measured bulk and mass-weighted mean values of that sample.

High-resolution δ^{13} C measurements on two carbonate SRMs suggest that kinetic isotope fractionation cannot exceed 1.8% in the RPO instrument. This agrees with intra-sample δ^{13} C trends of the environmental samples analyzed for this study, which display a large range in δ^{13} C spread between fractions and are consistent with independently measured carbon source composition. Additionally, selected samples display δ^{13} C trends with temperature opposite of that predicted by kinetic fractionation. These results are strong evidence that RPO kinetic fractionation is small and that blank and mass-balance corrected δ^{13} C values of each CO₂ fraction reflect carbon source isotope composition to within 1–2‰.

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SUPPLEMENTARY MATERIAL

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