Temperatures of aqueous alteration and evidence for methane generation on the parent bodies of the CM chondrites

Weifu Guo *, John M. Eiler

Division of Geological and Planetary Sciences, California Institute of Technology, Pasadena, CA 91125, USA

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Abstract

Aqueous alteration of primitive meteorites was among the earliest geological processes during the evolution of our solar system. 'Clumped-isotope' thermometry of carbonates in the CM chondrites, Cold Bokkeveld, Murray, and Murchison, demonstrates that they underwent aqueous alteration at 20–71 °C from a fluid with $\delta^{18}O_{VSMOW}$ of 2.0‰ to 8.1‰ and $\delta^{17}O_{VSMOW}$ of −0.1‰ to 3.0‰. The $\delta^{13}C_{VPDB}$ values of these carbonates exhibit a negative correlation with the $\delta^{18}O_{VSMOW}$ of their formation waters, consistent with formation and escape of $^{13}$C-depleted CH$_4$ during aqueous alteration. Methane generation under these conditions implies that the alteration fluid was characterized by an Eh of $<0.67$ and pH $>12.5$ (or lower at the highest alteration temperatures). Our findings suggest that methane generation may have been a widespread consequence of planetesimal and planetary aqueous alteration, perhaps explaining the occurrence of methane on Titan, Triton, Pluto, and other Kuiper-belt objects.

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1. INTRODUCTION

Aqueous alteration occurred on the parent bodies of the carbonaceous chondrites within the first tens of million of years of solar system history (Endress et al., 1996). The conditions of these processes are poorly constrained, both because direct samples of reactant fluids are rarely, if ever, preserved (Zolensky et al., 2004), and because the altered solids consist of complex, fine-grained mixtures of phases that present challenges to equilibrium thermodynamic approaches to calculate temperature, oxygen fugacity, and other relevant variables. Previous estimates of the temperatures of CM chondrite alteration range from <20 to <170 °C (Keil, 2000), based on the stabilities of constituent phases (e.g., tochilinite; Zolensky, 1984) or oxygen isotope fractionations between carbonate and phyllosilicate in the matrix (Clayton and Mayeda, 1984). This first approach yields only upper temperature limits; the second is only valid if matrix carbonate and phyllosilicate achieved oxygen-isotope exchange equilibrium. Differences in $\Delta^{17}O$ between carbonate and matrix in these samples (Benedix et al., 2003) indicate this assumption is not valid. In any event, uncertainties regarding the reduced partition coefficient ratios of the relevant phyllosilicate phases (Sheppard and Gilg, 1996) engender large uncertainties in apparent temperatures based on this approach. Baker et al. (2002) recently determined the oxygen isotopic composition of the structurally bound water released from phyllosilicates in Murchison and suggested an alteration temperature of ~80 °C for the CM chondrites based on the oxygen isotopic fractionation between this water and matrix carbonate. However, this estimation also assumes mutual isotopic equilibrium between these two reservoirs of oxygen.

We used the carbonate clumped-isotope thermometer (Ghosh et al., 2006; Schauble et al., 2006) to determine the temperatures of carbonate precipitation in the CM chondrites. The carbonate clumped-isotope thermometer is based on a thermodynamic equilibrium that orders $^{13}$C and $^{18}$O into bonds with each other within the carbonate lattice, and is independent of the isotopic composition of any co-existing phase. Moreover, this temperature information, combined with the known temperature-dependence of carbonate-water oxygen isotope fractionation (Kim and O’Neil, 1997), allows us to determine the oxygen isotope composition of waters from which these carbonates grew.
2. SAMPLES AND METHOD

2.1. Samples

Carbonate in the CM chondrites is mostly calcite—one of the bases of previous calibrations of the carbonate clumped isotope thermometer (Ghosh et al., 2006; Ghosh et al., 2007)—and is relatively homogeneous in composition and texture (Brearley and Jones, 1998). We analyzed carbonates from 7 splits of three different CM chondrites: Cold Bokkeveld (3 splits), Murray (1 split), and Murchison (3 splits). Each split was prepared from a separate fragment of a whole rock meteorite sample and ground individually. These samples represent near-extremes to the range of extent of aqueous alteration of CM chondrites: Murchison and Murray retain ca. ~93% and ~87%, respectively, of their original coarse, anhydrous silicate (chondrules and CAIs), whereas Cold Bokkeveld retains only ~45% of this original coarse anhydrous material (i.e., its texture more closely approaches that of the CI chondrites) (Browning et al., 1996).

2.2. Carbonate clumped isotope thermometry

\(^{13}\)C and \(^{18}\)O in thermodynamically equilibrated carbonate minerals preferentially group or 'clump' together into the same carbonate ion group to form \(^{13}\)C\(^{18}\)O\(^{16}\)O\(^{16}\)O\(^2\) as opposed to being randomly dispersed. This preferential clumping can be expressed through the equilibrium constant for the exchange reaction, \(^{13}\)C\(^{16}\)O\(^{16}\)O\(^2\) + \(^{15}\)C\(^{16}\)O\(^{16}\)O\(^2\) = \(^{15}\)C\(^{16}\)O\(^{16}\)O\(^2\) + \(^{13}\)C\(^{16}\)O\(^{16}\)O\(^2\). Because the doubly-substituted isotope of the carbonate ion (\(^{13}\)C\(^{16}\)O\(^{16}\)O\(^2\)) has a lower zero point energy than its normal and singly-substituted relatives (\(^{12}\)C\(^{16}\)O\(^{16}\)O\(^2\), \(^{12}\)C\(^{16}\)O\(^{16}\)O\(^2\), and \(^{12}\)C\(^{16}\)O\(^{16}\)O\(^2\)), the above reaction is thermodynamically driven to the right, producing abundances of \(^{13}\)C\(^{16}\)O\(^{16}\)O\(^2\) higher than that expected for a random distribution of isotopes. This abundance excess varies as a function of the temperature of carbonate growth or equilibration and forms the basis of a geothermometer (Ghosh et al., 2006; Schauble et al., 2006). The extent of \(^{13}\)C\(^{16}\)O clumping in a carbonate mineral is determined through the measurements of anomalous enrichment of mass 47 (mainly \(^{13}\)C\(^{16}\)O\(^{16}\)O) in the CO2 derived from phosphoric acid digestion of that mineral (Ghosh et al., 2006). We define the mass 47 anomaly, \(\Delta_{47}\), as the difference between the measured value of \(R_{47}^\text{measured} = ([\text{mass } 47]/[\text{mass } 44])\) and the value of \(R_{47}^\text{expected}\) expected in that sample if its C and O isotopes are randomly distributed among all isotope groups: \(\Delta_{47} = (R_{47}^\text{measured} - 1) \times 1000\) (Eiler and Schauble, 2004).

2.3. Analytical method

2.3.1. Sample digestion

Each sample split consisted of a whole rock fragment weighing between 40 and 250 mg. Each split was ground to a grain size of \(<500 \mu\text{m} with an agate mortar and pestle and then reacted with anhydrous phosphoric acid at 25 °C for 18–24 h to extract CO2 from calcite (Table 1). Carbonates other than calcite (e.g., dolomite) are rare in CM chondrites (Brearley and Jones, 1998) and do not react actively with phosphoric acid at 25 °C (less than 20% reaction of dolomite within 24 h, compared to 100% reaction of calcite; e.g., Al-Aasm et al., 1990). Therefore cross-contamination from other carbonate components in CM chondrite should be negligible during our extraction.

We examined the contribution of CO2 from oxidation of organic matter by conducting stepped acid extractions at 25 °C (i.e., anticipating that early steps would be dominated by carbonate and later steps by oxidation of organics). CO2 evolved at an average rate of \(\sim76.2\) mmol/mg/day during the first 24 h, and after that at an average rate of \(\sim0.9\) mmol/mg/day, suggesting that CO2 extracted for the first 24 h contained a negligible contribution from the oxidation of organic matter. We did not attempt to characterize the isotopic compositions of CO2 fractions collected after the first 24 h in this study because their sizes were too small for \(\Delta_{47}\) analysis and their \(^{13}\)C, \(^{15}\)O and \(\Delta_{47}\) values are well known from previous stepped-extraction studies (Grady et al., 1988; Benedix et al., 2003).

The yield of CO2 from each split varied between 39.3 and 127.6 µmol/g. We observe that smaller splits consistently produce higher yields of CO2 than larger splits (95.5–127.6 µmol/g for 40.5–60.6 mg splits vs. 39.3–61.8 µmol/g for 195.3–253.4 mg splits; Table 1). A similar correlation between CO2 yield and split size was observed in a previous study of CM chondrites (Murchison splits in Grady et al., 1988), and could be due to either heterogeneity among sample splits (i.e., if carbonate is generally present as widely-separated ‘nuggets’, smaller splits will produce greater maximum CO2 yields) or that phosphoric acid digestion more effectively attacks carbonate in smaller samples (e.g., if large amounts of ground sample are relatively impermeable).

Because of the relatively low abundances of carbonates in CM chondrites and the rare nature of the samples, we generally obtained less CO2 from our samples (3.9–12.1 µmol) than analyzed in previous studies of \(\Delta_{47}\) of CO2 produced by acid digestion of carbonate (≤50 µmol; e.g., Ghosh et al., 2006).

2.3.2. Purification of sample CO2

CO2 produced by phosphoric acid digestion of CM chondrites is exceptionally rich in sulfur and organic contaminants that must be removed prior to isotopic analysis. Therefore each sample of extracted CO2 was purified by exposure to dry Ag3PO4 (to remove sulfur contaminants, e.g., H2S), passage through a series of glass traps cooled with ethanol slush (−78 °C) and pentane slush (−130 °C), and passage through a Supelco Q-Plot GC column held at −20 °C (mainly to remove hydrocarbon contaminants; Affek and Eiler, 2006). Samples sized before and after dry Ag3PO4 treatment suggest that sulfur contaminants constitute 19–62% of the sample gas derived from phosphoric acid digestion of CM chondrite splits (38–62% for 3 Murchison splits, 58% for 1 Murray split and 19–29% for 3 Cold Bokkeveld splits). The potential for remaining contaminants was monitored by analyses of masses 48 and 49 in
the purified CO₂ gas, because these ion beams are less intense than that for mass 47 and can be strongly elevated in gases containing volatile organic contaminants. Previous studies have shown that such contaminants can lead to correlations between relatively small mass 47 excesses and proportionately greater excesses in masses 48 and 49 (Eiler and Schauble, 2004). While we never succeeded in completely removing masses 48 and 49 contaminants from these samples, no such correlations between Δ⁴⁷ values and either mass 48 or 49 excesses were observed in these purified samples, suggesting that, in this case, the remaining contaminants are not associated with a mass-47 isobar. Consequently, no correction on measured Δ⁴⁷ was made to account for masses 48 and 49 enrichments.

Although the sample purification procedures used in this study resemble those used in previous studies involving Δ⁴⁷ measurements (e.g., Eiler and Schauble, 2004; Affek and Eiler, 2006; Ghosh et al., 2006), we apply them to considerably smaller samples. We examined the effect of sample size on analytical fractionations associated with cryogenic purification, sulfide removal on silver phosphate, gas chromatography and mass spectrometric measurements. We observe a systematic effect of sample size on measured Δ⁴⁷ value only for the gas chromatography step. In particular, Δ⁴⁷ values of small samples (less than ca. 15 μmol) are systematically higher (by up to 0.2‰) in Δ⁴⁷ value as compared to large (i.e., larger than 15 μmol) samples of that same gas subjected to the same gas chromatographic purification procedure. The magnitude of this Δ⁴⁷ increase in small samples is independent of the isotopic compositions of the CO₂ gas and varies only with the size of the gas (i.e., the smaller the size of CO₂ gas, the bigger is its Δ⁴⁷ increase after GC purification). It is possible that this sample-size effect reflects interaction between CO₂ and adsorbed water in the GC column (although the column is routinely heated to 150 °C between samples and the observed Δ⁴⁷ increases are independent of the starting Δ⁴⁷ values—inconsistent with the effects of CO₂–H₂O exchange). It is unlikely that the effect reflects fractionation associated with partial loss of the sample because we routinely achieved >95% sample recovery of CO₂. Nevertheless, despite the fact that the cause of this effect is unclear, it is a consistently and clearly observed artifact of GC purification. Therefore, we routinely standardized analyses of samples smaller than 15 μmol by comparison with standard CO₂ gases of the same size that had been heated to 1000 °C to achieve the stochastic distribution. We tested the validity of this standardization method by measuring ~5 μmol samples of CO₂ prepared by acid digestion of small aliquots of carbonate standards having known Δ⁴⁷ values (NBS-19 and Sigma carbonate; Ghosh et al., 2006). Values of Δ⁴⁷ determined for these small aliquots of NBS-19 (n = 4) and Sigma carbonate (n = 4), standardized to heated gas of the same size, average 0.36 ± 0.03 (1σ, one standard deviation) and 0.56 ± 0.05 (1σ), respectively, consistent with their established values (0.35 ± 0.02, 0.55 ± 0.03; Ghosh et al., 2006). Therefore we believe that small samples can be analyzed with no systematic error provided they are compared to reference gases of the same size processed in the same way. Note however, that while developing these methods for analyses of small samples, some ini-

<table>
<thead>
<tr>
<th>Sample</th>
<th>Murchison</th>
<th>Cold Bokkeveld</th>
<th>Cold Bokkeveld</th>
<th>Estimated external precision (1σ)</th>
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<td>12/23/05</td>
<td>12/24/06</td>
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<td>61.8</td>
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<tr>
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<tr>
<td>δ¹⁸O (‰/VPMDB)</td>
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<td>δ¹⁷O (‰/VSMOW)</td>
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<td>34.1</td>
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<td>δ⁴⁴C (‰/VSMOW)</td>
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<td>δ¹⁷O (‰/VSMOW)</td>
<td>20.65</td>
<td>20.65</td>
<td>20.65</td>
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</tr>
</tbody>
</table>

a MAI—Mineralogical Alteration Index (Browning et al., 1996), with z = 0.35, 64 and are used in O¹⁷ corrections of this study.
tial tests of Sigma carbonate yielded CO₂ having Δ_{47} values consistent with room-temperature equilibrium of CO₂ gas (~0.98‰). We believe this reflects the great susceptibility of small samples to exchange with water adsorbed on surfaces in the gas-handling apparatus, necessitating unusually aggressive measures to dry the apparatus between samples (i.e., relatively long baking).

It is possible that exposure of CO₂ gas to dry Ag₃PO₄ may also influence the Δ_{47} of CO₂ samples. To evaluate this possibility, we prepared clean CO₂ gas and divided it into 2 splits of approximately equal size. We passed one split only through the GC column and the other split was first exposed to dry Ag₃PO₄ and then passed through the GC column. Values of Δ_{47} for these two splits were within 2 standard errors of each other, suggesting that exposure to dry Ag₃PO₄ does not influence the Δ_{47} value. Purifications on meteorite samples were performed only after such tests showed that a given batch of Ag₃PO₄ had negligible influences on Δ_{47} values.

2.3.3. Mass spectrometric analysis

Mass spectrometric analyses of purified CO₂ were performed in dual inlet mode on a Finnigan-MAT 253 gas source isotope ratio mass spectrometer, configured to simultaneously measure masses 44, 45, 46, 47, 48, and 49. Detailed descriptions of the mass spectrometer configuration are given by Eiler and Schauble (2004) and Ghosh et al. (2006). Two types of dual inlet measurements were employed in the present study: (1) the 3 analyses made in 2005 were performed at signal intensities of 2 V for mass 44 in regular dual inlet mode; (2) the 4 analyses made in 2006 were performed at signal intensities of 3.4–7.4 V for mass 44 in microvolume dual inlet mode (i.e., purified sample CO₂ was cryogenically concentrated in a small volume during measurements, instead of being expanded to the sample bellow). Measurements of each sample gas consist of 6–10 analyses, each of which involves 10 cycles of sample-standard comparison with an ion integration time of 8 s per cycle. Pressure balance was maintained at mass 44 between sample gas and reference gas for all analyses. Values of Δ_{47} for each sample were established by comparison with the same type of analyses of CO₂ gases of similar size and bulk composition that had been heated to 1000 °C (establishing the Δ_{47} = 0‰ reference scale for that sample size and purification procedure; Ghosh et al., 2006).

It has been previously shown that CO₂ derived from acid digestion of CM chondrite carbonates differs in oxygen isotope composition from the terrestrial mass fractionation line (Clayton and Mayeda, 1984; Benedix et al., 2003). We must consider this fact when performing ion correction calculations associated with our measurements. To compute the δ^{13}C and δ^{18}O values for these samples from their measured masses 44, 45, and 46 abundances, we assumed the average carbonate Δ^{17}O (recalculated from Benedix et al., 2003) for samples from the same CM chondrite and a value of λ = 0.5164 for the terrestrial mass fractionation line (Santrock et al., 1985). Values of δ^{13}C and δ^{18}O were standardized by comparison with CO₂ generated by phosphoric acid digestion of NBS-19 and are reported vs. VPDB and VSMOW, respectively.

Carbonate formation temperatures were estimated from the Δ_{47}–T calibration line for calcite and aragonite determined in Ghosh et al. (2006): Δ_{47} = \frac{-0.0034+0.02}{T} – 0.02, where T is in Kelvin. Note that Δ^{17}O values also affect calculated Δ_{47} values since the Δ_{47} value depends on δ^{13}C, δ^{18}O, and δ^{16}O values, and thus one must know or assume a Δ^{17}O value when calculating Δ_{47} (Affek and Eiler, 2006). For example, a 1‰ decrease in Δ^{17}O, if unrecognized or unaccounted for, would lead to a systematic error of 0.034‰ in Δ_{47}. The Δ^{17}O values of carbonates in any given CM chondrite vary by 0.1–0.8‰ (Murchison, n = 5; Murray, n = 2; Clayton and Mayeda, 1984; Benedix et al., 2003), thus introducing ~1 to ~5 °C uncertainty in our temperature estimation.

Because our measurements were made on relatively small samples and required extensive purifications of CO₂ produced by phosphoric acid digestion, they are less precise than previously reported temperatures based on carbonate clumped isotope thermometry (ca. ±10 °C 1σ vs. ±2 to 3 °C 1σ). External precisions of individual measurements of δ^{13}C, δ^{18}O, and Δ_{47} were estimated based on analyses of ~5 μmol samples of CO₂ prepared from small aliquots of carbonate standards and purified in the same manner as gases derived from CM chondrite samples. External precisions for these standard analyses averaged 0.14‰ for δ^{13}C, 0.21‰ for δ^{18}O and 0.041‰ for Δ_{47} (all 1σ; see Section 2.3.2). Similarly, analyses of clean ~5 μmol CO₂ gases exposed to dry Ag₃PO₄ and/or GC purification demonstrated external precisions averaging 0.05‰, 0.12‰, and 0.039‰, respectively (all 1σ). We assume the larger of these two sets of external precisions (those for small aliquots of carbonate standards) apply to our measurements of CM chondrite carbonates.

3. RESULTS AND DISCUSSION

3.1. Temperatures of aqueous alteration and isotopic compositions of the alteration fluid

Six out of the seven temperatures of carbonate formation determined using the carbonate clumped isotope thermometer are within the range of 20 to 35 °C—indistinguishable from each other given our analytical precision (Table 1). The exception is a split from Cold Bokkeveld, which yielded a significantly higher temperature of 71 °C. The similarity in temperatures of aqueous alteration among most splits of Cold Bokkeveld, Murray, and Murchison suggests that they reacted with water at similar depths in and stages of evolution of their respective parent bodies. This result is broadly consistent with previous thermal models for the parent bodies of the carbonaceous chondrites, which suggest that aqueous alteration occurred within a relatively narrow range of locations within their host parent bodies (Young, 2001; McSween et al., 2002). Note, however, that the higher temperature we observe for one split of Cold Bokkeveld indicates that aqueous alteration occurred over a range of temperatures in at least one sample.

If we interpret our results in the context of the model of Young (2001) that describes the thermal evolution of small planetesimals, water–rock reaction could have occurred within a ~250 m thick zone, 1.8 km from the surface of
an assumed 9 km radius parent body, although the higher temperature observed in one split requires that alteration occurred in a region, perhaps somewhat deeper, that was heated to well above the melting point of water ice. This compares with the estimate of an alteration zone 100 m thick, 1 km from the surface previously inferred based on oxygen isotope geochemistry and the mineralogical alteration index (Benedix et al., 2003).

Our measurements of the $\delta^{13}$C and $\delta^{18}$O values of CM chondrite carbonates (in particular the calcite fraction extracted by our acid digestion methods) vary from 37.77‰ to 68.88‰ and from 28.21‰ to 34.61‰, respectively (Table 1). This variation is similar to what has previously been observed for carbonates in CM chondrites ($\delta^{13}$C = 23.7‰ to 67.8‰; $\delta^{18}$O = 20.0‰ to 37.2‰; Clayton and Mayeda, 1984; Grady et al., 1988; Brearley et al., 1999; Benedix et al., 2003). However, the $\delta^{13}$C values of our samples of Murchison carbonates differ markedly from those in Grady et al. (1988) (60.09–68.88‰). An inverse correlation exists between $\delta^{13}$C and $\delta^{18}$O of Murchison carbonates considered in this study, in contrast to the weak positive correlation observed for eleven splits of Murchison in Grady et al. (1988). We suspect this difference reflects the fact that CM chondrites are highly heterogeneous breccias and contain alteration phases that may have formed over a range of conditions; thus, it is unsurprising that repeated sampling of small aliquots of such materials can produce divergent results.

Given the temperatures of carbonate growth summarized above and previous experimental calibration of the oxygen isotope fractionation between calcite and water (Kim and O’Neil, 1997, corrected for the phosphoric acid digestion factor used in this study $a = 1.01025$ at 25°C; Sharma and Clayton, 1965), we can estimate the oxygen isotope compositions of waters from which the carbonates we analyzed grew. These waters varied in $\delta^{18}$OVMOW between values of 2.0‰ and 8.1‰ (±2‰, 1σ). If we assume that the carbonates we analyzed have $\Delta^{18}$O values equal to the averages previously determined by Benedix et al. (2003), our results also indicate that the $\delta^{13}$O values of waters from which carbonates grew varied between –0.1‰ and 3.0‰. These $\delta^{18}$O and $\delta^{13}$O values of carbonate formation water are significantly lower than those estimated for the primary source of water that altered the CM chondrites ($\delta^{18}$O ca. $\geq 15.9$‰, $\delta^{13}$O ca. $\geq 9.2$‰; Clayton and Mayeda, 1999). This difference in isotopic composition between model reactant water and the water from which carbonates grew suggests that water evolves toward lower $\delta^{18}$O, $\delta^{17}$O, and $\Delta^{18}$O as a result of reaction with host rock (Clayton and Mayeda, 1984; Clayton and Mayeda, 1999) (Fig. 1).

We interpret our results in the context of a quantitative model for the aqueous alteration of CM chondrites. This model simplifies many aspects of this process, but provides a framework for testing the internal consistency of our interpretations. First, we assume, following Clayton and Mayeda (1999) and Eiler and Kitchen (2004), that the most volumetrically important alteration reactions (i.e., silicate hydration) can be represented by:

$$\text{Mg}_2\text{Si}_4\text{O}_{10} + \text{MgSiO}_3 + 2\text{H}_2\text{O} \rightarrow \text{Mg}_3\text{Si}_2\text{O}_5(\text{OH})_4$$

(1)

We define the reaction progress parameter (Clayton and Mayeda, 1999), $p$, as $p = \frac{x}{\gamma}$, where $\gamma$ is the fraction of initial olivine and pyroxene that has been converted to serpentine (on a molar oxygen basis) and $x$ is the ratio of moles of O in reactant water to moles of O in reactant olivine and pyroxene (thus $p$ denotes the ratio of moles of oxygen in the reacted rock to moles of oxygen in initial water). We assume that the initial oxygen isotopic composition of reactant water (i.e., prior to any reaction with anhydrous silicate) was $\delta^{18}$O$_{w,i} = 28.1$‰, $\delta^{13}$O$_{w,i} = 17.7$‰, and that the oxygen isotopic composition of reactant silicate was $\delta^{18}$O$_{s,i} = -4.2$‰, $\delta^{13}$O$_{s,i} = -7.4$‰ (Clayton and Mayeda, 1999).

Most of our model calculations adopt a serpentine–water oxygen isotope fractionation of $\Delta^{18}$O$_{w} = 12.5$‰ (Wenner and Taylor, 1971), appropriate for equilibration at 28°C. This corresponds to a $17$O/16O fractionation between serpentine and water of $\Delta^{17}$O$_{w} = 6.4$‰ (assuming a normal mass-dependent fractionation between serpentine and water). We selected a model temperature of 28°C because it is a reasonable approximation of the temperatures of carbonate growth for most of our samples (though silicate hydration need not occur at the same temperatures as carbonate precipitation) and it is similar to temperatures of aqueous alteration assumed by previous models of this kind (e.g., Clayton and Mayeda, 1999). While we are aware of no definitive reports of serpentinization occurring at 28°C in terrestrial rocks, both lab experiments and field observations have been used to suggest that serpentinization (and, important for our further discussion of this model in Section 3.2 of this paper, concurrent H$_2$ generation) occurs at earth-surface temperatures (Neal and Stanger, 1983; Stevens and McKinley, 1995; Stevens and McKinley,
The lowest measured temperature of a serpentinization reaction is 45°C, in the Lost City hydrothermal field, determined using a hydrogen isotope geothermometer (Proskurowski et al., 2006). However, the authors of that study suggested that this low temperature reflects re-equilibration of hydrogen isotopes after serpentinization at 110–150°C. Experimental studies suggest that hydration reactions are generally geologically fast (e.g., at 245°C with a rock/water ratio of 2.5/1, ~28% enstatite converts to serpentine within 6 days; Martin and Fyfe, 1970) and, in most circumstances, the rate limiting step is transport of water to the reaction site rather than the temperature-dependent kinetics of the reaction itself (Martin and Fyfe, 1970). We conclude that it is plausible to imagine sericite hydration on the CM parent bodies occurring at temperatures as low as 28°C, although a more definitive understanding of this problem will require further experimental studies of the kinetics of sericite hydration at such low temperatures. We also present a version of this model that assumes serpentine–water oxygen isotope fractionations more appropriate for a higher temperature of 71°C—the highest carbonate precipitation temperature observed in this study (Δ18Ow,w = 8.5‰, and Δ17Ow,w = 4.4‰; Wenner and Taylor, 1971) (Fig. 1).

Given the parameters summarized above, the oxygen isotope composition of aqueous fluid after reaction is given by Clayton and Mayeda (1999):

\[
\delta^{18}O_{\text{w,f}} = \frac{\delta^{18}O_{\text{w,i}} + (\delta^{18}O_{\text{s,i}} - \frac{1}{2} \Delta^{18}O_{\text{w,w}}) \rho}{1 + \rho} \quad (1)
\]

\[
\delta^{17}O_{\text{w,f}} = \frac{\delta^{17}O_{\text{w,i}} + (\delta^{17}O_{\text{s,i}} - \frac{1}{2} \Delta^{17}O_{\text{w,w}}) \rho}{1 + \rho} \quad (2)
\]

\[
\Delta^{17}O_{\text{w,f}} = \left( \frac{1 + \delta^{17}O_{\text{w,f}}/1000}{(1 + \delta^{18}O_{\text{w,f}}/1000)^{0.5\rho}} - 1 \right) \times 1000 \quad (3)
\]

Note that Δ17Ow,f values are independent of alteration temperature and vary only as a function of ρ once the initial isotopic compositions of water and rock (i.e., Δ17Ow,i and Δ17Ow,i) are specified. Therefore Δ17Ow,f (i.e., Δ17Ocarbonate, assuming carbonate formed in isotopic equilibrium with its formation water) can be used as an indicator for the extent of alteration. In contrast, the slope of the correlation between δ18Ow,f and δ17Ow,f is influenced by the alteration temperature (the higher the alteration temperature, the steeper is the line of δ18Ow,f vs. δ17Ow,f).

Our results imply that the water that precipitated carbonate in Cold Bokkeveld varied significantly in oxygen isotope composition over the course of that rock’s alteration history (Table 1). In the model trend depicting the expected isotopic evolution of reactant water illustrated in Fig. 1 (i.e., the higher extent of alteration the lower δ18O of the water), this water had experienced less previous reaction when the precipitation temperature was high and more previous reaction when the precipitation temperature was low; i.e., the rock was cooling as alteration proceeded. Previous studies of the minor and trace elements (Ricupiti et al., 1994) and stable isotope geochemistry (Grady et al., 1988) of carbonates in CM chondrites have also suggested that their parental waters varied in composition both spatially and temporally.

### 3.2. Correlation between δ13C of the carbonates and δ18O of their formation waters: evidence for methane generation

The δ13C values of carbonates in the CM chondrites exhibit a negative correlation with the oxygen isotope compositions of the waters from which they grew (Fig. 2). If we interpret this correlation in the context of the water evolution trend marked in Fig. 1, it suggests that the pool of dissolved inorganic carbon from which carbonate grew evolved from lower δ13C to higher δ18O as alteration proceeded. This interpretation is supported by weak correlations previously observed between the alteration index and both oxygen and carbon isotope compositions of CM chondrite carbonates (Grady et al., 1988), which was interpreted by the authors as possible indications that isotopically distinct components were altered and added to the fluid phase over the course of aqueous alteration.

It is possible that the trend in Fig. 2 reflects oxidation and dissolution of 13C-enriched carbon in the reactant assemblage. Carbon reservoirs, other than carbonates, in CM chondrites include organics (δ13CVPDB of ca. −36‰ to 41‰ for the major constituents; Sephton and Gilmour, 2001) and presolar grains (δ13CVPDB of bulk measurements on some presolar phases vary up to ca. 1500‰; Yang and Epstein, 1984). Organic constituents are lower in δ13C than the highest values observed in CM chondrite carbonates and so they cannot be the sole contributors to observed carbon isotope variations of those carbonates. Most presolar phases are highly resistant to chemical oxidation and dissolution (Yang and Epstein, 1984) and so should have been
inert under the relatively mild conditions at which CM chondrites underwent aqueous alteration. Presolar diamond and graphite might be more vulnerable to aqueous alteration, but their average $\delta^{13}C$ is also lower than that of CM chondrite carbonates (Hoppe et al., 1995). Thus, this hypothesis seems implausible.

Alternatively, $^{13}C$ enrichment could have accompanied aqueous alteration if it occurred in an open system (Wilson et al., 1999) from which CO$_2$ gas was lost by a mechanism that involved a kinetic isotope fractionation. Such processes lead to $^{13}C$ enrichment of dissolved inorganic carbon in evaporating Dead Sea brines (Stiller et al., 1985), caliches (Knauth et al., 2003) and cryogenic carbonates (Clark and Lauriol, 1992). However, significant CO$_2$ degassing in evaporating Dead Sea brines is caused by low pH of those brines (3.7–6.0; Stiller et al., 1985). Alteration fluids that reacted with the CM chondrites are inferred to have had far higher pH values (10–12; Zolensky, 1984) and are not expected to have evolved a substantial fraction of their dissolved inorganic carbon as CO$_2$. More importantly, kinetic isotope effects associated with evaporation of carbonate-bearing aqueous solutions typically also lead to increases in the $\delta^{10}O$ value of the residual solution (Stiller et al., 1985; Knauth et al., 2003). There is no evidence for such $^{18}O$ enrichment in the solutions that altered the CM chondrites—in fact, the opposite trend is observed in Fig. 2. Therefore, this process also seems like a poor explanation of our observations.

Finally, aqueous alteration of the CM parent body could have led to an increase in $\delta^{13}C$ of dissolved inorganic carbon in the pore waters of that body if alteration was accompanied by production and escape of low-$\delta^{13}C$ CH$_4$. A terrestrial analogue for this process is serpentinization of ultramafic rocks, during which dissolved inorganic carbon species are converted to CH$_4$ by the reaction (Horita and Berndt, 1999):

$$HCO_3^- + 4H_2 \rightarrow CH_4 + OH^- + 2H_2O \quad \text{(II)}$$

Palmer and Drummond (1986) and Shock (1988) suggest that reactions like reaction (II) require a condensed phase catalyst to be geologically significant. Chromite, Fe–Ni alloys, and magnetite have been shown to catalyze this reaction (Horita and Berndt, 1999; Foustoukos and Seyfried, 2004) and these and similar phases are abundant constituents of the CM chondrites (Becraft and Jones, 1998). At 200 °C (a temperature at which this reaction occurs over laboratory time scales), the CH$_4$ produced by this reaction is ca. 60% lower in $\delta^{13}C$ than the residual dissolved inorganic carbon (Horita and Berndt, 1999). Based on the temperature dependence of similar reactions (Horita, 2001), we suggest the fractionation at 28 °C (assuming the reaction could proceed over geological timescales at such low temperatures) could be as large as 90%. In this case, the observed range in $\delta^{13}C$ values for CM chondrite carbonates could be produced by ca. 38% loss of the dissolved inorganic carbon pool through reduction to and loss of CH$_4$. If we instead assume the experimentally measured high-temperature fractionation of 60%, then 51% loss would be required.

The carbon isotope composition of dissolved inorganic carbon varies with the progress of reaction (II), by the relation:

$$\delta^{13}C_{HCO_3^-} = 1000 \ln \left( \frac{a_{HCO_3^-}}{a_{HCO_3^-}} \right) = 1000 \ln \left( \frac{a_{HCO_3^-}}{a_{HCO_3^-}} \right)$$

$$\approx 1000 \left( \frac{F}{F_{13C_{CH_4,HCO_3^-}}} \right)$$

$$= 1000 \ln \left( \frac{F}{F_{13C_{CH_4,HCO_3^-}}} \right)$$

where $F$ is the fraction of reactant HCO$_3^-$ that remains and $x^{13C_{CH_4,HCO_3^-}}$ is the kinetic carbon isotope fractionation associated with reaction (II) (i.e., 90‰).

We prescribed that the progress of reaction (II) is a function of the progress of reaction (I), which is reasonable because H$_2$ is a reactant in reaction (II) and is produced in Fe-bearing systems as a result of reactions like reaction (I) (e.g., through olivine oxidation; Oze and Sharma, 2005, or Fe metal oxidation; Zolotov and Shock, 2004). This leads to:

$$F = \frac{[HCO_3^-]}{[HCO_3^-]} = 1 - \left( \frac{d[HCO_3^-]}{[HCO_3^-]} \right)$$

$$= 1 - \left( \frac{K[HCO_3^-]p}{[HCO_3^-]} \right)$$

where $[HCO_3^-]$ is the concentration of HCO$_3^-$ and $K$ is the ratio of the progress of reactions (I) and (II). Eq. (5) simplifies to: $F = e^{-\Delta}$. Thus, both the $\delta^{13}C$ of inorganic carbon are functions of $p$, and the position of a sample along the trend in Fig. 2 depends only on $p$ and $K$. If we assume that the initial $\delta^{13}C$ value of dissolved inorganic carbon was 22.8‰, the lowest $\delta^{13}C$ value observed for carbonates in the CM chondrites (23.7‰; Grady et al., 1988) minus the carbon isotope fractionation between dissolved inorganic carbon and calcite (~0.9‰; Rubinson and Clayton, 1969), then the trend in Fig. 2 is best fit by a value for $K = \frac{\Delta}{\Delta + 1}$ of 0.66, with $p$ varying between 0.72 to 1.19 among samples. Because $p$ denotes the ratio of moles of oxygen in the reacted rock to moles of oxygen in initial water, this variation of $p$ among samples indicates that water in some samples underwent ~1.6 times reaction with silicates than water in other samples.

Our model also predicts an inverse correlation between $\delta^{17}O$ and $\Delta^{17}O$ of the formation water on one hand and $\delta^{13}C$ of the carbonate on the other (Fig. 3). Most of the available measurements of $\delta^{13}C$ (Grady et al., 1988) and $\Delta^{17}O$ (Clayton and Mayeda, 1984; Benedict et al., 2003) in carbonates from the CM chondrites were made on separate sample aliquots and thus do not clearly test this prediction (although the ranges of their variations are broadly consistent with our predicted trend; Fig. 3). The only exception is a recent study on carbonates in paired Antarctic CM chondrites (Tyra et al., 2007), which observed an inverse correlation between $\delta^{13}C$ and $\Delta^{17}O$, as predicted. However, this study also observed that both $\delta^{13}C$ and $\Delta^{17}O$ correlate with $^{13}C$ content in the carbonates, and thus the correlation between $\delta^{13}C$ and $\Delta^{17}O$ could be caused or modified by terrestrial weathering (i.e., mixing or replacement of primary carbonates with lower $\delta^{13}C$, higher $\Delta^{17}O$ and $^{13}C$ terrestrial carbonates) (Tyra et al., 2007).

The correlation between $\Delta^{17}O_{water}$ and $\delta^{13}C_{carbonate}$ predicted by our model (Fig. 3) implies that carbonates should
vary in $\delta^{17}$O between different splits of the same meteorite for both Murchison and Cold Bokkeveld (i.e., because they vary in $\delta^{13}$C; Table 1). If our model is correct, then our assumption that $\delta^{17}$O values of carbonates from a given meteorite are uniform and equal to the previously measured average for carbonates from that meteorite introduces errors in $\Delta_{7}$ (which is calculated based on a known or assumed $\Delta^{17}$O value) and $\delta^{18}$O$_{\text{water}}$ (which depends, in part, on $\Delta_{7}$). These errors might potentially influence the correlation in Fig. 2, which is the primary justification of our methane generation model. A sensitivity analysis suggests that this factor is unlikely to lead to large changes: based on observed $\delta^{13}$C$_{\text{carbonate}}$ values and our model trend in Fig. 3, values of $\delta^{17}$O in carbonates are expected to vary by 0.3‰ or less among different splits from the same meteorite; this would result in variations of only 0.01‰ in ion-corrected $\Delta_{7}$ values. These variations would be within our nominal analytical uncertainties. It is possible to calculate an alternate version of Fig. 2 that assumes the model trend in Fig. 3 is correct. However, this is an essentially circular exercise (i.e., because the model trend in Fig. 3 depends, in part, on the fit to data in Fig. 2) and cannot clarify the issue. We suggest that future studies should attempt to apply both $\Delta_{7}$ and $\delta^{18}$O analyses to aliquots of CO$_2$ produced by acid digestion of carbonates in each CM chondrite.

3.3. Eh, pH of the alteration fluid estimated through the GEM-Selektor program

Reaction (II) will proceed to the right only under specific Eh conditions. We estimated these conditions using the GEM-Selektor program (Kulik et al., 1997; Kulik et al., 2000), which determines the equilibrium species distribution in a system through Gibbs energy minimization (Bethke, 1996). This method does not depend on the pre-alteration mineralogy (an uncertainty in previous models of this kind; Zolensky et al., 1989; Rosenberg et al., 2001).

Our model of the chemistry of fluids in equilibrium with the alteration assemblage in the CM chondrites considers Si, Fe, Mg, S, and Ca and neglects Al, Cr, K, Ni, and other minor elements. Calculations were performed for 28 °C, 1 atm, unless stated otherwise, with a bulk chemical composition as estimated by Browning and Bourcier (1998), assuming that aqueous alteration occurred without gain or loss of major cations (Table 2). During the simulation, we monitored the redox condition of the system by adding variable amounts of H$_2$ (0–1 g) into the system (which typically consisted of ~100 g of rock and 36 g of water, and thus corresponds to $p = 1.023$) and observed the concurrent changes of carbon speciation.

Controls were imposed on the final (post-alteration) mineral products to suppress the precipitation of mineral phases that generally do not form at these low temperatures (e.g., andradite, tremolite, and troilite etc.) or are kinetically inhibited (e.g., quartz). Chrysotile is the only

Table 2
GEM (Gibbs Energy Minimization) simulations of aqueous alteration on CM chondrite parent body

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>28</th>
<th>28</th>
<th>28</th>
<th>28</th>
<th>28</th>
<th>71</th>
</tr>
</thead>
<tbody>
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<td>Pressure (atm)</td>
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<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Initial unaltered bulk composition</td>
<td>Rock$^a$ (g)</td>
<td>94.91</td>
<td>94.91</td>
<td>94.91</td>
<td>94.91</td>
<td>94.91</td>
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<tr>
<td></td>
<td>Water (g)</td>
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<td>36</td>
<td>36</td>
<td>36</td>
<td>36</td>
</tr>
<tr>
<td></td>
<td>p$^a$</td>
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<td>0.5</td>
<td>2</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>CO$_2$ (M)</td>
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<td>0.1</td>
<td>0.1</td>
<td>0.01</td>
<td>1</td>
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<tr>
<td>Simulation results</td>
<td>Eh$^b$</td>
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<td>-0.66</td>
<td>-0.69</td>
<td>-0.67</td>
<td>-0.67</td>
</tr>
<tr>
<td></td>
<td>pH$^b$</td>
<td>12.5</td>
<td>12.3</td>
<td>12.7</td>
<td>12.5</td>
<td>12.4</td>
</tr>
</tbody>
</table>

$^a$ Rock here denotes a bulk composition typical of the fine-grained materials in CM chondrites (Browning and Bourcier, 1998), i.e., 31.15% SiO$_2$, 30.54% FeO, 21.89% MgO, 9.32% S, 2.01% CaO, and 5.09% others (wt%). “5.09% others” are not included in the simulations; $p$ is defined as the ratio of moles of oxygen in the reacted rock to moles of oxygen in initial water and is calculated based on the masses of rock and water given in the table; CO$_2$ denotes the initial concentration of inorganic carbon present in the pre-alteration assemblage, in the unit of mol/L (water).

$^b$ Eh and pH here denotes the Eh and pH conditions during the coexistence of calcite and CH$_4$ determined through GEM simulations (see text for simulation details).
Mg-phyllosilicate allowed to form in the simulation, consistent with the fact that it is the dominant Mg-rich alteration phase observed in carbonaceous chondrites (Brearley and Jones, 1998; Rosenberg et al., 2001). A typical product assemblage in our simulation included: chrysotile, greenalite, pyrite, pyrrhotite, and CH₄ (more reducing conditions); chrysotile, greenalite, pyrite, goethite, gypsum, and calcite (less reducing conditions). All the thermodynamic data employed in the calculations are from the Nagra/PSI Chemical Thermodynamic Data Base 01/01 (Hummel et al., 2002) and an updated version of thermodynamic database (slop98.dat) for SUPCRT92 (Johnson et al., 1997, 2000).

Given these constraints, we found that the coexistence of carbonate and CH₄ requires an Eh of −0.67 and pH of 12.5. Both Eh and pH are inversely correlated with the assumed temperature (decreasing to −0.73 and 11.7 at 71 °C), but neither of them is strongly dependent on reasonable variations of other model parameters (e.g., water/rock ratios, alteration pressures, and initial CO₂ concentrations etc., Table 2). For a given set of independent constraints, pH of the alteration fluid is inversely correlated with Eh. Note that this calculation (as for similar previous calculations; Zolensky et al., 1989; Rosenberg et al., 2001) assumes equilibrium with respect to redox reactions in the alteration assemblage, which might not apply (Bethke, 1996).

4. IMPLICATIONS

The temperatures of carbonate precipitation in representative CM chondrites, as constrained by carbonate clumped isotope thermometry, are within the range of (though more specific than) previous estimates based on mineral stability and oxygen isotope fractionation between different matrix phases. We observe a negative correlation between δ¹³C_PDB of carbonates and δ¹⁸O_VSMOW of the water from which those carbonates grew. This correlation can be explained if carbonate growth was accompanied by the formation and escape of ¹³C-depleted methane during aqueous alteration on the CM chondrite parent bodies.

Our conclusion that aqueous alteration of the CM chondrites was accompanied by formation and loss of methane implies that methane generation in the interiors of planets and planetesimals may have been a widespread phenomenon in the evolutionary history of the solar system. Several recent studies (Formisano et al., 2004; Lyons et al., 2005; Oze and Sharma, 2005) invoked serpentinization and hydrothermal alteration as sources of atmospheric methane detected on Mars, and analogous processes could be considered to explain methane found on other solar system objects, such as Titan (Niemann et al., 2005), Triton (Cruikshank et al., 1993), Pluto (Owen et al., 1993) and other Kuiper belt objects (Brown et al., 1997, 2005; Barucci et al., 2005; Licandro et al., 2006). It is unclear whether the details of the alteration model we propose (i.e., reactants, temperatures, Eh, pH, and stoichiometries) could have any direct relevance for Mars, which has undergone extensive magmatic differentiation and thus consists of relatively volatile-poor igneous rocks that differ in many respects from the carbonaceous chondrites. However, the model we suggest for methane generation in the CM chondrite parent bodies plausibly applies to at least some portions of the interiors of Titan and other outer solar system bodies, especially early during their histories. For example, it is possible that the subsurface of Titan contains abundant liquid water-ammonia solution with temperatures up to 0 °C (Grasset and Pargamin, 2005) and possibly higher early in Titan’s history (Atreya et al., 2006; Tobie et al., 2006). This reduced, water-rich interior provides an environment where silicate alteration reactions such as those we consider in our model could take place. Currently methane in the atmosphere of Titan is destroyed through photochemical reactions at a rate of 2.5 × 10⁻¹⁴ mol cm⁻² s⁻¹ (Yung et al., 1984). If Titan’s atmosphere is in steady state, this loss could be supported by methane produced by serpentinization of silicates in the presence of carbonate-bearing fluids at a rate of ~3.2 × 10¹² kg of silicate consumed per year (Oze and Sharma, 2005). If this rate is typical of the history of Titan, it would lead to time-integrated hydration of a ~39 km-thick layer of reactant rock. Similarly, the methane on Pluto is hydro dynamically escaping with a flux of 2.2 ± 10¹⁰ mol cm⁻² s⁻¹ (Trafton et al., 1997). This flux could have been supported throughout the history of Pluto by gradual aqueous alteration of a 34 to 157 km-thick global layer of reactant silicate rock. In both cases, the layer of reactant rock could be several times thinner if it is rich in metallic iron (because metallic iron is a more efficient source of H₂ during aqueous alteration than Fe(II)-silicates; Zolotov and Shock, 2004; Oze and Sharma, 2005).

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