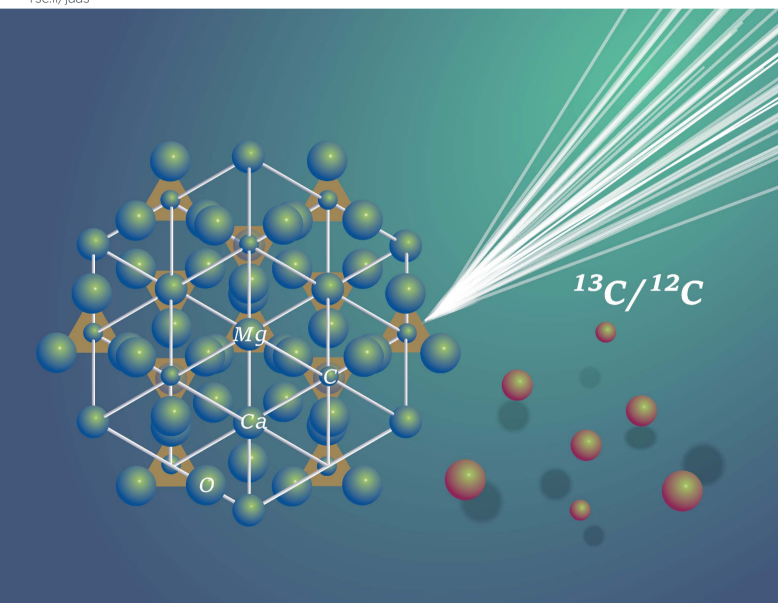
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Determination of carbon isotopes in carbonates (calcite, dolomite, magnesite, and siderite) by femtosecond laser ablation multi-collector ICP-MS†

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This study reports a method for in situ determination of stable carbon ($\delta^{13}C_{\infty}$) isotope compositions for calcite, dolomite, magnesite and siderite by femtosecond laser ablation multi-collector inductively coupled plasma mass spectrometry (fs-LA-MC-ICP-MS). The chemical and bulk C isotope compositions of various carbonates have been characterized by LA-ICP-MS and isotope ratio mass spectrometry (IRMS), respectively. Magnesite shows limited chemical variation, whereas the major element compositions of dolomite and siderite vary greatly among samples and individual grains. The spectral interference of doubly charged ions (24 Mg $^{2+}$ for 12 C $^+$ and 26 Mg $^{2+}$ for 13 C $^+$) and polyatomic species (12CH+ for 13C+) was evaluated. The associated internal precision of in situ C isotope analysis correlates with intensities and is better than 0.20% (2SE) when the measured 12 C ion signal is >12.5 volts. Carbonate samples DOL-8, MGS-1 and SD-5 show relatively uniform bulk and in situ carbon isotope compositions with precisions better than 0.24% and 0.45% (2SD) and are adopted as in-house standards. The laser ablation analysis results of various calcite, dolomite, magnesite and siderite samples with δ^{13} C in the range of -6.28% to 0.32% agree excellently with IRMS determined values using standard-sample bracketing (SSB) calibration, with a precision of 0.37-0.68% (2SD). Matrix effects have been investigated and determined to be insignificant for the same carbonate matrix with variable chemical compositions when adopting a femtosecond laser. Instrumental carbon isotope fractionation observed among different carbonate matrices is non-negligible (e.g., up to 4.29% between calcite and magnesite) using fs-LA-MC-ICP-MS, and an external multi-reference calibration can be applied for nonmatrix matched analyses.

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1. Introduction

Carbon occurs as both a major and trace element in variable amounts in various Earth systems (atmosphere, hydrosphere, and lithosphere). It has three isotopes, including two stable isotopes of 12 C (98.89%) and 13 C (1.11%) and a radioactive isotope 14 C ($\sim 10^{-12}$). The significant relative mass difference ($\sim 89\%$) between 13 C and 12 C results in a wide range of C isotope compositions in nature. By far, carbonates are the most abundant carbon-bearing minerals in the Earth's crust, with calcite

and dolomite $(CaMg(CO_3)_2)$ accounting for at least 90% of crustal carbon in massive sedimentary and metamorphic formations.² Other abundant carbonates including magnesite, siderite and rhodochrosite occur widely in sedimentary rocks, and in numerous metalliferous ore deposits in magmatic, hydrothermal, or sedimentary environments.³ Carbon isotope analyses of a variety of carbonates contribute to the understanding of terrestrial geochemical cycles given the number of natural processes that result in a large variation of δ^{13} C values between approximately -100% and +20%.^{4,5}

Conventional isotope ratio mass spectrometry (IRMS) allows for accurate and highly precise stable carbon isotope measurements of the absolute abundance of stable carbon isotopes in natural or synthetic materials. Recently, the reported analytical precision (sample-to-sample, 1SD) of $\delta^{13}C$ values measured for carbonate consuming 30 μg of sample was better than 0.04% (1SD) using a ThermoTM MAT253 coupled to a ThermoTM Kiel IV carbonate device. Nevertheless, the interpretation of C isotopic signatures acquired by conventional

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methods (*i.e.*, IRMS) is limited since they represent averaged (*i.e.*, multi-grain) isotopic values for individual samples.

In situ C isotopic analyses by secondary ionization mass spectrometry (SIMS),7-12 NanoSIMS13 and laser ablation coupled to multi-collector inductively coupled plasma mass spectrometry (LA-MC-ICP-MS)14 have received increased interest in Earth science-related studies, which provides valuable isotopic information of geological samples at a high resolution. For example, single carbonate crystallites (dolomite-ankerite) as small as 3- $10 \mu m$ across can be readily analyzed by SIMS with a sub per-mil (%) accuracy and precision (2s).10 The micron-scale C isotope signatures of magmatic calcium carbonate (calcite) have been successfully measured using ns-LA-MC-ICP-MS with a satisfactory precision (within 0.50%; 2SD).14 Due to the complex chemical and physical features of natural carbonate, efforts have been made to calibrate a series of dolomite-ankerite solid solution reference materials for C isotope microanalysis using SIMS.11 For example, a compositional bias was recorded in C isotope values for dolomite, which increased exponentially by 2.5-5.5% with increasing Fe contents.11 Without exception, instrumental bias resulting from compositional dependency (i.e., matrix effect) remains as one of the analytical challenges for the wide application of in situ C isotope analyses for various carbonate matrices using LA-MC-ICP-MS. Chen et al. (2017) successfully used nanosecond laser-Nu plasma MC-ICP-MS to determine the carbon isotope of calcium carbonate, 14 whereas the development of the carbon isotope microanalytical method of calc-magnesian, magnesian and ferruginous carbonates still remains a challenge. It is challenged by the lack of matrixmatched standards and C isotope fractionation at both the laser ablation site and ICP site.

In this last regard, femtosecond (fs) laser ablation systems are characterized by a much shorter pulse duration, even shorter than the thermal relaxation time of materials compared to the more popular nanosecond laser, which avoids heating laser plasma interaction. 15,16 Moreover, femtosecond lasers generate smaller and more uniform distribution of aerosol particles (usually on the nanoscale), which enhances the transmission efficiency and ion signal intensity.15-21 Less thermal effects and generation of small particles result in high ion signal intensity and reduced fractionation, which makes femtosecond laser ablation systems most effective in reducing matrix effects for in situ isotopic analyses by the LA-MC-ICP-MS method. 15-17,20 For example, accurate copper isotope compositions of cuprite samples were successfully determined using a pure Cu metal standard by fs-laser ablation analysis. 18 Isotopic fractionation induced by the matrix dependency of Fe bearing minerals when calibrated against a pure metal isotopic standard was not observed using fs-LA-MC-ICP-MS.22 In this study, we adopt a femtosecond laser to reduce carbon isotope fractionation at the laser ablation site for the analysis of calcmagnesian, magnesian and ferruginous carbonates. Here, we characterize a series of in-house carbonate reference materials and evaluate the feasibility of in situ C isotopic analysis for carbonates (calcite, dolomite, magnesite, and siderite) by fs-LA-MC-ICP-MS.

Experimental

2.1. In situ chemical composition analyses

The major and trace element concentrations for all investigated carbonate samples were determined with a RESOlution 193 nm laser system attached to an iCAP-Q ICP-MS at the State Key Laboratory of Geological Processes and Mineral Resources (GPMR), China University of Geosciences in Wuhan. NIST SRM 612 and USGS reference glass materials (BIR-1G, BCR-2G and BHVO-2G) were repeatedly analyzed for each 7–10 unknowns for calibration, as proposed by Chen *et al.* (2011). NIST 612 was used to correct for instrument drift and the USGS glass materials were used for external calibration for concentration determination. Both standards and samples were measured using a 33 μ m spot size, a 10 Hz repetition rate, and a corresponding energy density of \sim 3 J cm $^{-2}$. Offline data reduction, including the calculation of composition, detection limits and individual run uncertainties were calculated with ICPMSDataCal software. 24

2.2. Bulk carbon isotope analyses

Bulk carbonate mineral separates were crushed and milled to <75 µm for carbon isotopic determination using a MAT253 isotope ratio mass spectrometer (IRMS) coupled to a Thermo Fisher Gasbench II interface at the State Key Laboratory of Biogeology and Environmental Geology (BGEG), China University of Geosciences (Wuhan). The powdered carbonate samples ($\sim100-500~\mu g$) were placed into a 10 mL borsyl headspace vial and reacted with 500 µL of 99% orthophosphoric acid (MERCK, Darmstadt, Germany, catalog #1.00565) for ~5 hours at 72 °C. The δ^{13} C values of the evolved CO₂ were measured and reported as per mil relative to the Vienna Pee Dee Belemnite (VPDB) reference.²⁵ The data quality was monitored *via* repeated analyses of the internal laboratory standard GBW04416 (δ^{13} C = +1.61%, which yielded an analytical precision of better than 0.1%, (1 σ).

2.3. In situ carbon isotope analyses

In situ C isotope analyses were performed on a 257 nm Yb fslaser ablation system (NWR-Femto^{UC}, USA) coupled to a NEPTUNE Plus MC-ICP-MS instrument (Thermo Fisher Scientific, Bremen, Germany) at GPMR. The duration of the laser pulse is 300 fs, the wavelength is 257 nm, the maximum energy is 120 mJ, and the maximum pulse repetition rate is 1000 Hz. 13C and 12C were measured simultaneously in Faraday cups H3 and L4, and the ²⁵Mg²⁺ ion signal was acquired in the central cup. The samples were ablated in a helium atmosphere (0.6-0.7 L min⁻¹) within the ablation cell, which was then mixed with argon gas (1.0-1.5 L min⁻¹) in the sample-out line downstream from the ablation chamber before entering the plasma torch. A signal-smoothing device was used downstream from the sample cell to decrease the tau effect, which helps eliminate the short-term variability of the signal.26 All measurements were performed in low mass resolution mode (M/M as defined by a peak edge width from 5 to 95% of the full peak height) of \sim 400, whereas medium mass resolution (\sim 4000) was adopted to resolve spectral interference. The details of the instrumental

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Table 1 Operating conditions for the MC-ICP-MS and laser ablation system

Instruments	nents Operating conditions					
MC-ICP-MS (Neptune plus)						
Cup-configuration	L4 ¹² C	$^{ m C}_{^{25} m Mg^{2+}}$	H3 ¹³ C			
RF power	1150–1200 W	<u> </u>				
Cool gas flow	16.0 L min ⁻¹					
Auxiliary gas flow	$1.0~\mathrm{L~min^{-1}}$					
Argon make-up gas flow	\sim 1.0–1.5 L min $^{-1}$					
Helium carrier gas flow	0.6-0.7 L min ⁻¹					
Nitrogen gas flow	$0~\mathrm{mL~min}^{-1}$					
Interface cones	X skimmer cone + standard sample cone					
Mass resolution	${\sim}400~\mathrm{(low)}$					
Block number	1					
Cycles of each block	120					
Integration time (s)	0.524 s					
Laser ablation system (NWR-FemtoU	JC, USA)					
Laser type	Yb:YAG femtosecond laser					
Wavelength	257 nm					
Pulse length	300 fs					
Energy density	$3.24 \ \mathrm{J} \ \mathrm{cm}^{-2}$					
Ablation mode	Raster mode					
Spot size	60 μm					
Raster scan rate	$10 \; \mu m \; s^{-1}$					
Laser frequency	80 Hz					

operating conditions and measurement parameters summarized in Table 1.

The offline selection and integration of background and analytical signals, and time drift correction and isotopic ratio calibration were performed using Iso-Compass software.²⁷ In situ C isotope data reported were corrected for instrumental mass bias using the standard-sample bracketing method. In order to allow for the comparison of the in situ δ^{13} C data determined here relative to those of in-house reference materials, all data presented were recalculated to the commercially available VPDB marine carbonate reference material25 using the following equations:

$$\delta^{13}C = \left[\frac{(^{13}C/^{12}C)_{\text{sample}}}{(^{13}C/^{12}C)_{\text{standard}}} - 1 \right] \times 1000 \tag{1}$$

$$\delta^{13}C_{sample/VPDB} = \delta^{13}C_{sample/in\text{-}house\ RM} + \delta^{13}C_{in\text{-}house\ RM/VPDB}(2)$$

where $\delta^{13}C_{\text{sample/in-house RM}}$ is the measured δ value of the sample relative to the in-house reference materials by fs-LA-MC-ICP-MS and $\delta^{13}C_{\text{in-house RM /VPDB}}$ is the δ value of the in-house reference materials relative to VPDB determined by IRMS.

3. Results and discussion

3.1. Chemical and bulk isotopic characterization of carbonates

Different types of isomorphic series of carbonate minerals were adopted in this study, including 6 dolomites (DOL-1, DOL-2, DOL-3, DOL-6, DOL-8, and DOL-9), 3 magnesites (MGS-1, MGS-2, and MGS-3) and 5 siderites (SD-1, SD-2, SD-3, SD-4,

and SD-5), together with two previously characterized calcites (SXD8 and Oka153).14

The major and trace element compositions of the carbonate materials are listed in Appendix Table S1† and shown in Fig. 1. The results of LA-ICP-MS analysis show that the major element composition of magnesite is relatively uniform with Mg contents that range between 26.3 and 27.4%, while that of dolomite and siderite varies greatly among samples and within individual grains. Dolomite displays variable Mg and Fe contents that range between 8.6-12.5% and 0.03-5.4%, respectively, whereas Mn and Ca show limited variation (abundances vary within 1.5%). Most dolomite grains are characterized by relatively homogeneous major element compositions (2SD < 0.6%), whereas DOL-1 and DOL-6 exhibit within-grain variations. For instance, the Fe and Mg contents of DOL-1 vary from 0.5-4.0% and 9.0-11.5%, respectively, and those of DOL-6 show ranges between 5.1-6.5% and 7.4-9.1%. The siderite samples show variable Mg contents (2.4-8.6%), whereas they in general contain little Mn and Ca (<1.8% and 0.2%, respectively). Among them, SD-4 contains the highest Mg (average of 8.6%) and the lowest Fe contents (average of 32.2%). Trace element contents for the investigated carbonates are mainly lower than 10 $\mu g g^{-1}$, with the exception of Na, Al and Sr that are several to hundreds of $\mu g g^{-1}$ (Appendix Table S1†).

Three to six randomly selected chips were taken from each carbonate sample to obtain the bulk carbon isotope values using IRMS (e.g., Fig. 2), and data are shown in Table 2. The bulk δ^{13} C values of the examined dolomite define a range of -5.27% to 0.32\%, and those of magnesite and siderite range between 0.16\% and 3.00% and -9.78% to -5.32%, respectively. Among the investigated carbonate samples, DOL-8, MGS-1 and SD-5 show

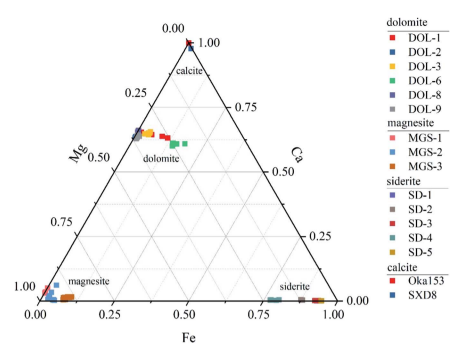


Fig. 1 Major-element variations of different carbonate matrices (dolomite, magnesite, siderite and calcite).

good reproducibility within 0.25% (2SD), suggesting relative isotopic homogeneity on the hand-specimen scale. The carbonates with larger determined uncertainties (e.g., >0.50\%, 2SD) show C isotopic variations for distinctive chips separated; for instance, different siderite crystals taken from the SD-2 sample are characterized with distinctive C isotope compositions as -4.90%, -5.20% and -5.87%, respectively (Fig. 2).

3.2. Spectral interference

For the determination of C isotopes using fs-LA-MC-ICP-MS, spectral interference involves both polyatomic species

(dominantly carbon hydroxide ion) and doubly charged Mg ions, and tetravalent charged Ti and Cr ions. Both the sample and carrier gases of argon and helium contain CO2 or organic impurities, which results in ¹²CH⁺ interference of ¹³C. The ¹²CH⁺ interference, which occurs on the high mass side of the ¹³C ion signal, can be significant and is dependent on both the Ar and He gas flow rates. Fig. 3 shows the influences of the Ar and He gas flow rates on CH⁺ interference examined in medium resolution mode (resolving power ~ 4000). $^{12}CH^{+}/^{13}C$ ratios increase from 0 to 0.333 when the Ar gas flow rate is augmented from 1.208 to 1.463 L min⁻¹ (Fig. 3c). Similarly, this interference

Table 2 fs-laser ablation C isotope measurements for the carbonate samples^a

Sample Ma		5 1 d	fs-LA values	fs-LA values			IRMS values		
	Matrix	Bracketing standard	δ^{13} C (‰)	2SD (‰)	n	δ^{13} C (‰)	2SD (‰)	n	
Oka153	Calcite	SXD8	-5.61	0.42	26	-5.77^{b}	0.04^b	1	
DOL-8	Dolomite	DOL-8	-1.06	0.37	22	-1.09	0.08	3	
DOL-9	Dolomite		0.41	0.65	28	0.32	0.30	3	
DOL-2	Dolomite		-5.61	0.59	35	-5.27	0.62	3	
DOL-3	Dolomite		-1.15	1.41	33	-0.92	1.58	3	
DOL-1	Dolomite		-2.64	3.19	41	-0.96	0.94	3	
DOL-6	Dolomite		-2.76	0.81	15	-2.14	0.83	3	
MGS-1	Magnesite	MGS-1	3.04	0.40	24	3.00	0.12	6	
MGS-2	Magnesite		0.49	0.57	16	0.16	0.51	6	
MGS-3	Magnesite		0.91	0.99	20	0.70	0.78	3	
SD-5	Siderite	SD-5	-6.26	0.45	22	-6.21	0.24	3	
SD-1	Siderite		-6.14	0.68	15	-6.08	0.16	3	
SD-2	Siderite		-6.37	0.54	13	-5.32	0.98	3	
SD-3	Siderite		-5.98	0.38	15	-6.28	0.08	3	
SD-4	Siderite		-10.11	0.98	18	-9.78	0.22	3	

^a "n" represents the number of laser ablation rastering and IRMS analyses, respectively. "2SD" represents twice the standard deviation. ^b Data from Chen et al. (2017).14



Fig. 2 Photograph of the hand specimen of SD-2 siderite with marked sampling positions of chips 1, 2, and 3 and associated IRMS determined C isotope compositions.

also becomes more significant with increasing He gas flow rates (Fig. 3d). The 12 CH $^+$ / 13 C interference can be avoided by carefully tuning and adopting low gas flow rates for both Ar and He (<1.208 and 0.6 L min $^{-1}$).

All investigated carbonates in this study are low in Ti and Cr (<10 $\mu g~g^{-1}$; Appendix Table S1†), and the tetravalent charged ion interference of $^{48}\text{Ti}^{4+}$ for $^{12}\text{C}^+$ and $^{52}\text{Cr}^{4+}$ for $^{13}\text{C}^+$ is negligible. In contrast, the spectral interference of doubly charged Mg ions may be significant for C isotope analyses of magnesium-rich carbonate minerals, such as dolomite and magnesite (Fig. 4). $^{26}\text{Mg}^{2+}$ and $^{24}\text{Mg}^{2+}$ interference both occur on the low mass shoulder of ^{13}C and ^{12}C ion signals and thus the latter was measured at an off-center position (high mass shoulder) and in interference-free plateau region (Fig. 4).

3.3. Internal precision

The internal uncertainty (or precision; repeatability within the ablation raster) for an individual fs-LA-MC-ICP-MS analysis ranged from 0.13 to 1.14‰ and varied according to C signal intensities; the latter was achieved by modifying laser ablation parameters such as an ablation spot size of 45–60 μm , scan rate of 2–12 μm s $^{-1}$, frequency of 25–80 Hz and fluence of 2.7–3.3 J

cm⁻² (Fig. 5). Fig. 5 clearly shows the dependency of the internal precision on the 12 C ion signal intensity. Of note, an increase in the 12 C ion signal intensity from 3 to 12.5 V results in a significant improvement of the repeatability, and ideally measured 12 C ion signals should be >12.5 V in order to achieve a level of precision <0.2% (2SE).

3.4. Reproducibility

The external reproducibility (level of duplication) associated with the in situ C isotope analysis of carbonates is in general influenced by instrument stability, reliability of mass bias correction, correction for spectral interference, and homogeneity of the C isotope composition of individual samples.28 The in situ reproducibility of carbonate samples with relatively good bulk carbon isotope reproducibility (including DOL-8, MGS-1, and SD-5) were further verified in this study by investigating individual carbonate grains. The dimensions of the three carbonate grains examined are approximately 1.5 cm \times 2 cm, 1 cm \times 1.5 cm and 1 cm \times 1 cm, respectively (e.g., Fig. 6a). The external reproducibility of the measured C isotope ratios was obtained from repeated measurements with the same experimental/instrument conditions. The associated

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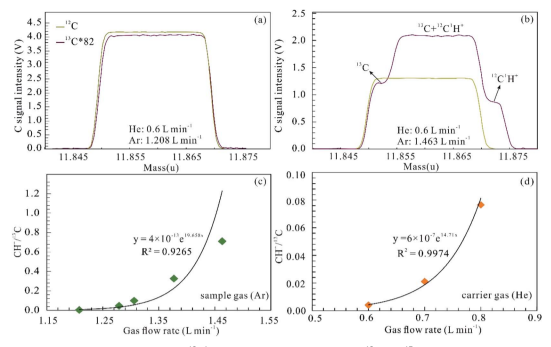


Fig. 3 Influences of the Ar and He gas flow rates on ${}^{12}C^{1}H^{+}$ interference. Peak shapes for ${}^{12}C$ and ${}^{13}C$ of the background at different Ar and He gas flow rates (a and b), and $^{12}C^1H^+$ superimposed on the high mass shoulder of ^{13}C in (b). Signal intensity of ^{13}C was multiplied by 82 to match the signal intensity of ¹²C. ¹²C¹H⁺ interference (¹²C¹H⁺/¹³C) displays an exponential function of both the Ar sample gas flow (c) and the He carrier gas flow (d)

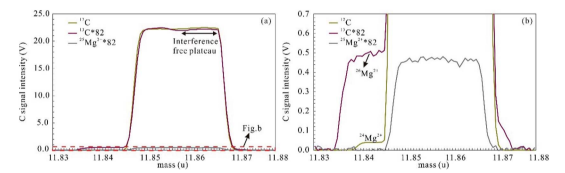


Fig. 4 Peak shapes for ¹²C and ¹³C obtained from MGS-1. Interference of doubly charged Mg ions superimposed on the low mass shoulder of ¹²C and 13 C, which leaves the high mass shoulder as the interference free plateau. Signal intensities of 13 C and 25 Mg²⁺ were multiplied by 82 to match the signal intensity of ¹²C.

uncertainty for each sample (external reproducibility) is expressed as twice the standard deviation (2SD). The within-run precision (2SE) of each analysis involves the uncertainty propagation from the bracketing standard materials, adopting the following equation as proposed in Zhang et al. (2020):27

$$2SE = 2 \times \sqrt{\left(SE_{sample}\right)^2 + \left(SE_{standard-1}\right)^2 + \left(SE_{standard-2}\right)^2}$$
(3

where standard-1 and standard-2 represent measurements of the bracketing standards performed before and after the sample, respectively.

Repeated analyses of C isotopes for different portions of the individual carbonate grains were performed by conducting

analyses along cross sections as shown in Fig. 6a. This yielded δ values with twice standard deviation of $-1.06 \pm 0.37\%$ (n = 22), $3.04 \pm 0.40\%$ (n = 24) and $-6.26 \pm 0.45\%$ (n = 22) for DOL-8, MGS-1 and SD-5, respectively (Fig. 6b). Repeated analyses of DOL-8, MGS-1 and SD-5 yielded δ^{13} C external reproducibility in the range of 0.37-0.45% (2SD), which are similar to those previously reported (~0.50%; 2SD) in Chen et al. (2017)14 using ns-LA-MC-ICP-MS, and therefore represent the resolving capacity of the in situ C isotope analysis protocol established in this study.

An alternatively preliminary homogeneity assessment has been conducted using the obtained data from different zones of the large investigated grain with the two-factor analysis of variance (ANOVA), according to ISO guide 35.29 The δ^{13} C results

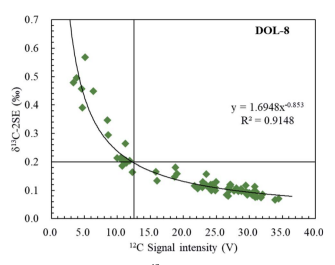


Fig. 5 Internal precisions of δ^{13} C (2SE) measured by femtosecond laser ablation MC-ICP-MS on DOL-8, as a function of signal intensity of 12 C (volts). It demonstrated that the measurement repeatability of the δ^{13} C ratio (SE) is an exponential function of the 12 C intensity.

of different zones in each carbonate were randomly separated into three units. F is the ratio of among-unit variance (S_{among}^2) to within-unit variance (S_{within}^2):

$$F = \frac{S_{\text{among}}^2}{S_{\text{within}}^2} \tag{4}$$

where $S_{\rm among}^2$ represents the ratio of the among-unit sums of squares (SS_{among}) to the associated among-unit degrees of freedom ($\nu_{\rm among}$), and $S_{\rm within}^2$ is the ratio of the within-unit sums of squares (SS_{within}) to the associated within-unit degrees of freedom ($\nu_{\rm within}$):

$$S_{\text{among}}^2 = \frac{SS_{\text{among}}}{v_{\text{among}}}$$
 (5)

Table 3 ANOVA table for homogeneity testing of carbon isotope compositions of magnesite, dolomite and siderite a

Sample	$ar{X}$	SS_{among}	SS_{within}	F	$F_{0.05}$ (2,30)
MGS-1	3.06	0.12	1.03	1.72	3.32
DOL-8	-1.10	0.10	0.92	1.54	3.32
SD-5	-6.24	0.09	1.64	0.78	3.32

$$S_{\text{within}}^2 = \frac{SS_{\text{within}}}{\nu_{\text{within}}} \tag{6}$$

where v_{among} and v_{within} are dependent on the number of units and the analysis number within each unit.

$$v_{\rm among} = m - 1 \tag{7}$$

$$v_{\text{within}} = N - m \tag{8}$$

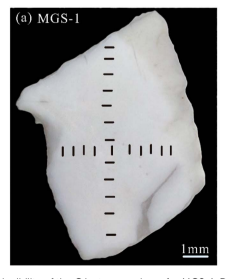
where m represents the number of units and N represents the total number of statistics.

The results are shown in the ANOVA table (Table 3). *F*-Testing demonstrates that the samples have very good C isotopic homogeneity $[F < F_{\text{critical}} \ (\nu_{\text{among}}, \ \nu_{\text{within}});$ the critical value of *F* for $\alpha = 5\%$].

To sum up, these three carbonate materials are considered to have isotopically homogeneous C isotope compositions on a μm to mm scale and are therefore adopted as in-house reference materials for carbon isotope analysis using fs-LA-MC-ICP-MS.

3.5. Results of *in situ* C isotope analysis of calcite, dolomite, magnesite and siderite using fs-LA-MC-ICP-MS

Multiple carbonate materials with a wide range of δ^{13} C (bulk sample) values ranging from -9.78 to 0.70% were investigated using the standard-sample bracketing strategy outlined here. The previously characterized SXD8 carbonate together with



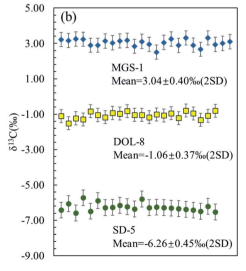


Fig. 6 External reproducibility of the C isotope analyses for MGS-1, DOL-8 and SD-5 (b) with the line scan ablation positions marked on MGS-1 (a).

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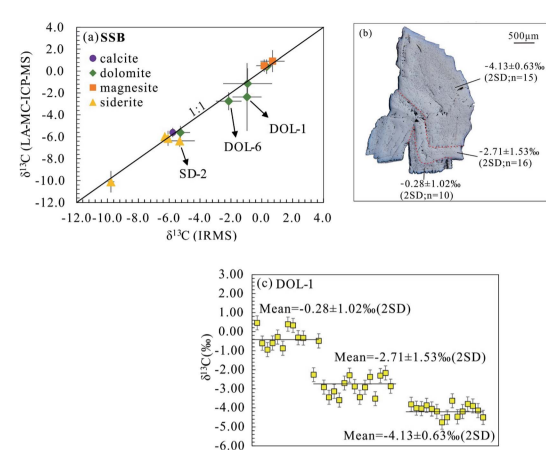


Fig. 7 Measured δ^{13} C values of dolomite, magnesite and siderite using fs-LA-MC-ICP-MS in comparison to IRMS data (a). The solid line is a 1 : 1 reference line. Laser ablation carbon isotope measurements of DOL-1 show three distinct *in situ* δ^{13} C values (b and c).

DOL-8, MGS-1 and SD-5 were used as the bracketing standards.¹⁴

Calcite. Both calcites (SXD8 and Oka153) are characterized by low Mg, Fe and Mn contents (<0.7%), and detailed chemical and isotopic information can be found in Chen *et al.* (2017). Separate calcite grains from sample Oka153 were analyzed using SXD8 as the bracketing standard yielding an average δ^{13} C value of $-5.61 \pm 0.42\%$ (2SD), which agrees well with that determined by IRMS (Table 2 and Fig. 7a). δ^{14}

Dolomite. The five dolomite samples selected for this investigation contain variable Mg and Fe contents (8.6-12.5% and 0.1-5.4%, respectively; Fig. 1 and Appendix Table S1†). Two dolomite samples (DOL-9 and DOL-2) with relatively homogeneous isotope ratios yield δ^{13} C values of 0.41 \pm 0.65% and $-5.61 \pm 0.59\%$, respectively (Table 2 and Fig. 7a). Both results are in good agreement with the bulk δ^{13} C analyse results obtained by IRMS. DOL-1 together with DOL-3 and DOL-6 displays a saddle texture (e.g., Fig. 7b), which is recognized as a common product of late-stage diagenesis and hydrothermal activity or as a replacement product in some dolomitized limestones. 30,31 The bulk carbon isotope compositions for these saddle dolomites DOL-1, DOL-3 and DOL-6 show slight heterogeneity as associated uncertainties are >0.83% (2SD), with in situ analyses yielding average δ^{13} C values of $-2.64 \pm 3.19\%$, $-1.15 \pm 1.41\%$, and $-2.76 \pm 0.81\%$ (2SD), respectively; despite their larger associated uncertainties, the average *in situ* C isotope values are consistent with those obtained for the corresponding bulk samples by IRMS. The recorded large uncertainty is indicated by distinct C isotope composition in different grains or zones. For instance, DOL-1 defines three distinct *in situ* δ^{13} C values of $-4.13\pm0.63\%_{o}$, $-2.71\pm1.53\%_{o}$, and $-0.28\pm1.02\%_{o}$ in different zones of the grain with generally heavier C isotopes identified in the rim (Fig. 7b and c). The recorded C isotopic variation indicates significant isotope fractionation during the diagenesis of the investigated saddle dolomites.

Magnesite. The magnesite crystals selected for this study are chemically homogeneous (Fig. 1). The reported *in situ* δ^{13} C value for MGS-2 using MGS-1 as the bracketing standard is 0.49 \pm 0.57% (2SD), which shows good agreement with IRMS measurements (Table 2 and Fig. 7a). The laser-ablation δ^{13} C analyses of MGS-3 display an average value and associated precision of 0.90 \pm 0.99% (2SD; Table 2 and Fig. 7a). The slightly larger uncertainty for MGS-3 compared to that of MGS-2 is consistent with that observed in the bulk C isotope analyses (Table 2), which implies that MGS-3 is isotopically heterogeneous on both the high-spatial micron scale and the hand-specimen scale.

Siderite. Average δ^{13} C values of $-6.14 \pm 0.68\%$, $-5.98 \pm 0.38\%$ and $-10.11 \pm 0.98\%$ were obtained, respectively, for samples SD-1, SD-3 and SD-4 by laser ablation analyses using

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SD-5 as the bracketing standard and show good agreement with the bulk sample IRMS values (Table 3 and Fig. 7a). Of particular note, *in situ* C isotopic results for SD-3 show excellent micronscale homogeneity (2SD = 0.38%0 and n=15) despite slightly variable chemical compositions (Table 2 and Fig. 1). Of note, laser ablation analyses for SD-2 yield an average δ^{13} C value and associated uncertainty of $-6.37 \pm 0.54\%$ 0 (2SD), which is slightly lower than the average IRMS value of $-5.32 \pm 0.98\%$ 0 (2SD), but similar to the same chip with IRMS determined δ^{13} C value of -5.87%0 (Fig. 2).

3.6. Carbon isotope fractionation among various carbonates

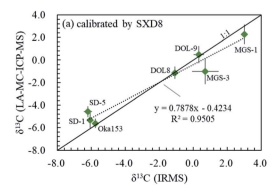
The range of carbon isotope signatures reported here for different carbonate minerals, which include calcite, dolomite, magnesite and siderite, using the fs-LA-MC-ICP-MS method is shown in Fig. 8. All C isotope compositions for carbonates investigated here are compared relative to calcite standards and were obtained during two analytical sessions on different analytical days. The possible matrix-dependent C isotope difference/fractionation between calcite (SXD8) and dolomite (DOL-8 and DOL-9) is variable and ranges from insignificant (first analytical session) to non-negligible (up to 2.89% in the second analytical session; Fig. 8a and b). The instrumental fractionation values between magnesite (MGS-1 and MGS-3) and siderite (SD-1 and SD-5) with respect to calcite are both non-negligible, up to 4.29% and 1.64%, respectively (Fig. 8a and b). The matrix difference can influence the accuracy by: (1) introducing isobaric interference and/or (2) altering the instrumental mass bias effected by plasma ionization and ion transmission efficiency, as well as (3) laser-induced fractionation at the site of ablation.32-35 Fe and Mg may be the dominant matrix elements within the investigated carbonate matrices inducing carbon isotope fractionation using fs-LA-MC-ICP-MS. As discussed above, the isobaric interference of doubly charged Mg ions is avoidable by conducting analyses on the interference free plateau, and thus it does not play the dominant role in the C isotope fractionation between calcite (SXD8) and Mg-bearing carbonate (e.g., dolomite DOL-8 and DOL-9 and magnesite MGS-1 and MGS-2). Therefore, the matrix difference dominantly results in a non-spectral

instrumental mass bias. The different sample physical and chemical properties of carbonate matrices could result in different aerosol particle sizes causing differential transport and vaporization efficiency, elemental/isotopic redistribution in aerosols and a hot melt layer due to diverse volatile properties of elements, different plasma temperatures and space charge by a distinct number of samples transmitted, all of which affect both plasma ionization and ion transmission efficiency and laser-induced fractionation.35 Evans et al. (2021)³⁶ reported that the matrix dependency of B isotope measurements is related to ICP-MS tuning conditions, especially RF power, and suggested that strict matrix matching does not appear necessary when using a session-specific correction line based on three well-characterized references. Similarly, Fig. 8 shows a good linear correlation between the LA results and IRMS data in both analytical sessions, and the external multi-reference calibration can serve as an alternative calibration approach in addition to the matrix-matched calibration. By adopting the linear correlation and the laser ablation C isotope results for multi-reference materials, accurate C isotope results (IRMS results in the formula) of carbonate samples can be calculated using the equation below:

$$\delta^{13}C_r = k \times \delta^{13}C_m + b \tag{9}$$

where $\delta^{13} C_{\rm m}$ is the LA analysis δ value, $C_{\rm r}$ represents the IRMS/true δ value, and k and b are the slope and the y-intercept of the linear regression and can be determined using multiple standards in routine analyses (Fig. 8a and b). The slope and y-intercept of the linear regression are affected by instrument conditions and operating engineers (Fig. 8a and b). This linear relationship should be established on a daily basis to guarantee the accuracy and precision of C isotopic analysis.

The instrumental mass bias for the measurement of C isotope ratios was also previously corrected using internal normalization adopting boron isotope in solution MC-ICP-MS.^{37,38} In their work, absolute carbon ¹³C/¹²C isotope amount ratios (corrected with ¹¹B/¹⁰B) have been obtained for four NIST reference materials with a precision better than 0.05%, which



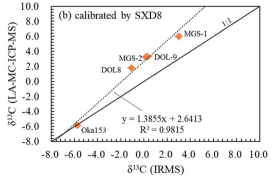


Fig. 8 Carbon isotopic fractionation of different carbonate minerals including calcite, dolomite, magnesite and siderite using fs-LA-MC-ICP-MS in two different analytical sessions (a and b). δ^{13} C obtained by fs-laser ablation is a function of that obtained by IRMS, represented by the dashed line, applying SXD8 calcite as a calibration standard. The solid line is a 1:1 reference line.

provides a possible option to correct instrument drift for the non-matrix matched calibration protocols. Detailed future work should be conducted in order to correct the non-spectral matrix effects for different carbonate matrices, especially rare minerals (e.g., stichtite and malachite) for which calibration standards have not yet been established.8

4. Conclusions

This study presents a method to obtain relatively high-precision in situ C isotope compositions for a variety of carbonates (calcite, dolomite, magnesite and siderite) by fs-LA-MC-ICP-MS. The internal precision of *in situ* C isotope analyses is better than 0.2% (2SE) for analyses with recorded ¹²C ion signal intensities >12.5 V, and the corresponding external reproducibility of inhouse standards DOL-8, MGS-1 and SD-5 is better than 0.45\% (2SD). Using the standard-sample bracketing (SSB) calibration protocol, the C isotopes of various carbonates with variable δ^{13} C signatures (-6.28-0.32%) are accurately determined with uncertainties that range between 0.37% and 0.68% (2SD). Moreover, in situ C isotope results obtained with the fs-LA-MC-ICP-MS protocol described here essentially eliminate matrix effects as they are relatively uniform for the same carbonate matrix with variable chemical compositions. Instrumental carbon isotope fractionation among the investigated different carbonate matrices (e.g., up to 4.29% between calcite and magnesite) are observed using fs-LA-MC-ICP-MS, and an external multi-reference calibration can be applied for nonmatrix matched analyses.

Conflicts of interest

There are no conflicts to declare.

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