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# Isotopic fractionation of oxygen and carbon in decomposed lower-mantle inclusions in diamond

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**Abstract** Two carbonatitic mineral assemblages, calcite + wollastonite and calcite + monticellite, which are encapsulated in two diamond grains from the Rio Soriso basin in the Juina area, Mato Grosso State, Brazil, were studied utilizing the NanoSIMS technique. The assemblages were formed as the result of the decomposition of the lower-mantle assemblage calcite + CaSi-perovskite + volatile during the course of the diamond ascent under pressure conditions from 15 to less than 0.8 GPa. The oxygen and carbon isotopic compositions of the studied minerals are inhomogeneous. They fractionated during the process of the decomposition of primary minerals to very varying values:  $\delta^{18}$ O from -3.3 % to +15.4 % SMOW and  $\delta^{13}$ C from -2.8 % to +9.3 % VPDB. These values significantly extend the mantle values for these elements in both isotopically-light and isotopically-heavy areas.

## Introduction

Over the last two decades, the mineralogical composition of the Earth's lower mantle has become known from mineral inclusions in diamond (e.g., Harte 2010; Kaminsky 2012; Harte and Hudson 2013 and references therein). However, the isotopic characteristics of lower-mantle mineral associations have been poorly studied to date. Only two ion-microprobe analyses from a grain of CaSi-perovskite within lower-mantle diamond have been performed; they have yielded values of  $\delta^{18}O$  at 4.92 % and 6.95 % SMOW (Harte et al. 1999). The former is close to the average mantle value, while the latter is shifted in the supracrustal values.

Our knowledge of isotopic 'average mantle values' of carbon and oxygen is mainly based on, and can be applied to, only the upper-mantle reservoir. The high-precision determination of the upper-mantle oxygen isotopic composition yielded the average value of  $\delta^{18}O = 5.3 \pm 0.6$  % SMOW (Valley et al. 1998; Spetsius et al. 2008; Wang et al. 2011; Kostrovitsky et al. 2012; Giuliani et al. 2014). The range of the upper-mantle carbon isotopic composition is from -2 to -8 %  $\delta^{13}C$  (VPDB, while for the lower mantle the range is suggested to be wider, from -0.5 to -8.5 %  $\delta^{13}C$  (Cartigny 2005). A detailed study of lower-mantle diamonds from Juina (Brazil) and Kankan (Guinea) confirmed the similarity of the lower-mantle and the upper-mantle carbon isotopic compositions (Palot et al. 2013).

We report here the first results of oxygen and carbon isotope analysis of other minerals identified in lower-mantle diamonds (Kaminsky et al. 2009; Wirth et al. 2009) utilizing the NanoSIMS technique.

## Samples and analytical method

This study investigated mineral inclusions encapsulated in two diamond grains from the Juina area, Mato Grosso State, Brazil. Both diamonds were extracted from the same prospecting

sample #8 collected within the Rio Soriso basin (see Fig. 2 in Kaminsky et al. 2009). These inclusions are daughter minerals, formed as a result of the decomposition of the initial carbonatitic association and form two assemblages: calcite + wollastonite II and calcite + monticellite (Fig. 1). Calcite and wollastonite II were analyzed within a  $4 \times 4$  µm rhombic-shaped inclusion in diamond #8/103, foil #1734 (Kaminsky et al. 2009). Calcite forms the matrix of the inclusion occurring as a polycrystalline aggregate with individual crystals on the order of 1-2 µm each (Fig. 1A). Its composition is essentially  $CaCO_3$  (Ca = 99.26 at. % of the cation group), with an admixture of Sr (0.74 at. % of the cation group) and trace quantities of Ba. Such Sr-enriched calcite is considered to be primary magmatic (e.g., Armstrong et al. 2004). Wollastonite II (high) was identified from its diffraction pattern; it occurs along with nyerereite and a Ca-garnet. The wollastonite II forms several near-euhedral grains, 50-500 nm in size. The chemical composition of wollastonite II is (in at. %): Ca = 49.7, Si = 50.1, and Fe = 0.2.

The calcite + monticellite assemblage was studied from diamond #8/104 (foil #1598) (Wirth et al. 2009). This plate-like inclusion, only a few micrometers in length and ca. 200 nm thick, is composed of calcite, that comprises the bulk of the inclusion, and sub-micrometre-sized monticellite, cuspidine, wollastonite II, metallic alloys, amorphous material with a C-Ca-Si-O composition, and minor fluid bubbles, 5–50 nm in size (Fig. 1B). The calcite, as in sample #8/103, is polycrystalline, with an average grain size of approximately 1–2  $\mu$ m. However, in this sample, the chemical composition of the mineral is that of pure calcite, that may be explained by its recrystallization. Monticellite (identified from its diffraction pattern) has a composition of (at. %): Si = 29.8, Fe = 11.4, Mg = 23.9, and Ca = 33.6, with minor admixtures of Al (0.7), Ti (0.3), and Mn (0.3).

Both diamonds belong to the same geological sample, ~0.5 m<sup>3</sup> in size. Both studied mineral assemblages in these diamonds belong to the natrocarbonatite association, in which nyerereite (Na,K)<sub>2</sub>Ca(CO<sub>3</sub>)<sub>2</sub> forms the matrix of the inclusion (Kaminsky et al. 2009, 2015).

The major challenge for isotopic study of these mineral grains was their very small, sub-micrometre, sizes. We performed the analysis of the samples utilizing a NanoSIMS 50 instrument at the Lawrence Livermore National Laboratory, USA. We measured the carbon and oxygen isotope compositions of minerals from the same FIB foils that had previously been examined by TEM at the GeoForschungsZentrum Potsdam, Germany (Fig. 1). Oxygen and carbon isotope abundances were determined using a focused primary 0.3 pA Cs<sup>+</sup> beam, ca. 60 nm in diameter, which was rastered over a 4 × 4 µm² area to produce quantitative secondary ion images. Each ion image was subdivided into calcite, silicate and diamond regions (Fig. 2), which were analyzed separately. An electron flood gun was used for charge compensation. Secondary ions were acquired by simultaneously measuring <sup>12</sup>C-, <sup>13</sup>C-, <sup>16</sup>O-, <sup>18</sup>O- and <sup>28</sup>Si- on electron multipliers at a mass-resolving power of ca. 3600. Isotope ratios were determined with the L'Image software developed by L. Nittler, Carnegie Institution of Washington, DC. One could expect the quasi-simultaneous arrival (QSA) effect on C isotopes (Slodzian et al. 2004). However, in our experiments, no variations of the instrumental fractionation factor with count rates were observed. Further an admixture of Sr (0.74 at. %) and a trace quantity of Ba are not so

high as to influence the O isotope composition (Valley and Kita 2009). Other elemental impurities (Mg, Fe, Mn) in the studied calcites are well below 1 at. %, so the results do not need matrix correction (Rollion-Bard and Marin-Carbonne 2011). Instrumental mass fractionation was assessed from two calcite standards and the UW-2 garnet standard; they cover the 22 ‰ of  $\delta^{13}C$  composition range. The external precision on the standards (84 hrs time) was < 4 ‰ for  $\delta^{18}O$  and < 9 ‰ for  $\delta^{13}C$  (2 $\sigma$ ).

## **Results**

The results obtained in our study were very surprising (Table 1). Against the expected mantle values, the  $\delta^{18}O$  values for the calcite + wollastonite II assemblage vary from -1.6 % to -3.3 %, and for the calcite + monticellite assemblage from +15.0 % to +15.4 % SMOW. The  $\delta^{13}C$  values for calcite are -2.8 % in foil #1734 (in association with wollastonite II) and +9.3 % VPDB in foil #1598 (in association with monticellite). Both carbonate-silicate assemblages are in isotopic equilibrium, having similar  $\delta^{18}O$  values for both mineral phases in each sample. At the same time, there are significant differences in the isotopic characteristics between the two samples:  $\Delta = 12.1$  % for carbon and  $\Delta = 16.6$ –18.7 % for oxygen.

Large error values (6.0–6.1 % for carbon and 4.3–8.2 % for oxygen) include not only analytical errors, but (mainly) isotopic inhomogeneity of samples at the nano-scale level, which is well seen in Fig. 2A,B, where isotopic variations in a ~2 µm calcite grain, reflected in counts per second, vary within ~3.000 ct/s for <sup>12</sup>C and ~100.000 ct/s for <sup>16</sup>O, that significantly exceeds the analytical errors. The isotopic heterogeneity of mantle minerals has been observed at both the centimeter- and micrometer-scale. For example, a study of a single diamond-bearing peridotite xenolith revealed that, within a volume of only 27 cm<sup>3</sup>, the variations for nitrogen content, nitrogen aggregation state, carbon and nitrogen isotopic compositions, respectively, cover 64 %, 75 %, 15 % and 23 % of the ranges known for all peridotitic diamonds (Thomassot et al. 2007). Carbon isotope compositional variations within a single diamond crystal reach 10 %  $\delta^{13}$ C and more (Kaminsky and Sobolev 1985). At the nano-scale the isotopic variations appeared to be even greater. Diamond #8/106 from Rio Soriso was studied for isotopic composition with NanoSIMS analysis at the Center for Advanced Marine Research, Ocean Research Institute at the University of Tokyo; fifteen spot measurements of  $\delta^{13}$ C in the diamond were found to vary from -14 % to -25 % VPDB, with an average at -19.40  $\pm$  3.22 % (Wirth et al. 2007). The isotopic study of  $\delta^{15}$ N in other diamond crystals from the Juina area in Brasil, performed with the use of a Cameca IMS ion microprobe at the Department of Terrestrial Magnetism, Carnegie Institution of Washington, USA, produced results with one-sigma values of  $\pm 3$  % (D. Araujo, pers. comm.). This means that the 'error values', in fact, are mainly ranges of isotopic heterogeneity for the studied samples. The real analytical errors are less than 0.4–1.7 ‰ (see Table 1 and Fig. 4); these values include as well, possible isotopic fractionation between

coexisting carbonate and silicate phases, which is particularly significant under low, subsolidus temperatures (100–500 °C) during the course of secondary alteration of the minerals (see below).

## Discussion and preliminary conclusions

In both analyzed samples calcite is the major mineral. Calcite in sample #1734 is enriched in Sr and, partly, in Ba. Such calcite is inferred to be primary-magmatic (e.g., Armstrong et al. 2004), and its  $\delta^{13}$ C value (-2.8 %) is close the range typical of mantle values. Its origin may be attributed to the lower mantle. Earlier, in diamond from the Rio Soriso area, calcite was found in a close ('touching') assemblage with walstromite-like CaSiO<sub>3</sub>, formed after decompressed CaSiperovskite, which is typical of the lower-mantle association (Brenker et al. 2007). One of the studied samples #8/103, in the same foil #1734, contains small (less than 100 nm), euhedral nano-inclusions of wüstite (Mg# = 1.9-15.3) in association with periclase (Mg# = 84.9–92.1) (Fig. 7 in Kaminsky et al., 2009). The coexistence of these phases implies their formation under pressures at or greater than 85 GPa and temperatures in excess of 1000 K (Dubrovinsky et al. 2001); i.e., within the lowermost part of the lower mantle. This conclusion is confirmed by the identification, in the same diamond, of the assemblage of iron carbides, Fe<sub>3</sub>C, Fe<sub>7</sub>C<sub>3</sub> and Fe<sub>23</sub>C<sub>6</sub> with native iron and graphite, which crystallized within a pressure interval of 50–130 GPa and, possibly, have their source in the outer core or within the D" layer of the lowermost part of the mantle (Kaminsky and Wirth 2011).

The other minerals, found in association with calcite, are stable at much lower pressures and temperatures. Wollastonite forms as a result of the decomposition of CaSi-perovskite, which starts at ca. 15 GPa with the formation of titanite (CaSi<sub>2</sub>O<sub>5</sub>), larnite (Ca<sub>2</sub>SiO<sub>4</sub>), walstromite (CaSiO<sub>3</sub>), and then, at ca. 4 GPa, wollastonite (CaSiO<sub>3</sub>) (Akaogi et al. 2004). Further uplift and resultant decompression to levels where pressures were less than 0.8 GPa initiated a reaction of wollastonite + fluid to form cuspidine + monticellite. This reaction should have occurred at comparatively low pressures because it is known, from experiments exceeding 0.8 GPa and 1450 °C, that monticellite breaks down to merwinite + forsterite (Kushiro and Yoder, 1964). Unfortunately, little is known about the pressure–temperature stability of cuspidine. Cuspidine was reported to occur in calc-silicate xenoliths from the Bushveld Complex, South Africa which had peak pressure–temperature conditions of 0.11–0.24 GPa and >1200 °C (Buick et al., 2000). The formation of cuspidine and monticellite is the latest-stage event during transit of the host diamond. The existing grains of wollastonite-II found as an inclusion in foil #1734 are metastable relics which were preserved in a closed system. After the consumption of fluorine, the formation of cuspidine at the expense of wollastonite stopped, and the present-day mineral assemblage is observed. The quantity of monticellite within the inclusion was limited by the availability of Al and Mg (Wirth et al., 2009).

Based on the data obtained, the major trends in the behavior of O and C isotopes during the decomposition of the primary lower-mantle mineral association may be preliminarily marked.

The  $\delta^{18}O$  SMOW values of the studied assemblages, compared with  $\delta^{18}O$  values of other diamond associations worldwide, are much dispersed, ranging from -3.3 ‰ to +15.4 ‰ (Fig. 3). They exceed not only values for the upper-mantle (from +4.7 to +5.9 ‰, after Valley et al., 1998), but all other analyses of primary mantle minerals. To our knowledge, there are no isotopic characteristics for carbonatitic fluid inclusions in diamond to date. (Klein-BenDavid et al. (2010) performed isotopic analyses for C, N, Sr, Nd and Pb from samples of fibrous diamonds, in which diamond matrix predominates.) Fig. 4 presents the  $\delta^{13}C$  vs.  $\delta^{18}O$  values for the investigated diamond inclusion assemblages, and compares these data to those from other mantle minerals and to the modified primary igneous carbonatite box after Srivastava et al. (2005). The samples, in their  $\delta^{13}C - \delta^{18}O$  coordinates, differ significantly from all previously studied kimberlitic and carbonatitic minerals. Two models can be offered to explain this.

The first model (A in Fig. 4) is an independent C and O fractionation for each sample from a single source. Sample #1598 has a very heavy oxygen isotopic composition, which, in other cases, for other mineral inclusions in diamond, has been explained by the involvement of crustal oxygen in the formation of diamond and associated minerals for the eclogitic association, as a result of deep lithospheric slab subduction (e.g., Jacob et al. 1994; Schulze et al. 2003, 2013; Ickert et al. 2013). However, the  $\delta^{13}$ C value in the same sample contradicts this suggestion: eclogitic-type diamonds are commonly characterized not by heavy, but, quite the reverse, by light carbon isotopic composition, from -10 % to -35  $\delta^{13}$ C VPDB. Only some eclogitic diamonds from New South Wales, Australia record values of -2 ‰ to +4 ‰ VPDB (e.g., Sobolev et al. 1989; Cartigny 2005). Such great  $\delta^{13}$ C values as those reported herein for sample #1598 (+9.3 to +15.4 % VPDB), have no analogues in igneous and metamorphic rocks. The value of  $\delta^{13}$ C in sample #1734 has another disagreement with known data on the isotopic systematics of diamond: while the  $\delta^{13}$ C values are within the average mantle range, the  $\delta^{18}$ O values (-1.6 % and -3.3 % SMOW) lie within a negative area, never before observed in diamond associations or terrestrial magmas (e.g., Criss 2008) for that matter. Such values are characteristic of only meteoric waters or metamorphic rocks that have been affected by such waters.

Suggesting a similar origin for both diamonds and their inclusions (belonging to the same geological sample), their C-O isotopic characteristics may be considered as the result of a single process, and belonging to different stages in the decomposition of the primary lower-mantle mineral association. In this case, an alternative model B (Fig. 4) may be suggested. The two samples should be considered as two ultimately differentiated values from an original single source, which had intermediate  $\delta^{13}C$  and  $\delta^{18}O$  values lying between the two samples.

The distinct compositions of calcite in the analyzed assemblages support model B. Calcite in sample #1734, associated with wollastonite II, is enriched in Sr and is primary-magmatic (e.g., Armstrong et al. 2004); its  $\delta^{13}$ C value (-2.8 ‰) lies within the range typical of mantle values. Calcite in sample #1598, associated with monticellite, is chemically pure and has a very heavy carbon isotopic composition,( $\delta^{13}$ C = +9.3 ‰). It may be suggested that, during the ascent of diamond to the surface, wollastonite + remaining fluid, encapsulated in diamond, was dissociated into monticellite + cuspidine (see above). Such dissociation occurs at low, subsolidus

temperatures (500 to 100 °C). According to the calculations performed by Wilson et al. (2007), the oxygen isotopic fractionation, in a closed kimberlitic system, within the temperature range from 1100 °C to 100 °C varies from 6 ‰ to 10 ‰. In this case, the  $\delta^{18}$ O values change from the average mantle values of +5 ‰ – +6 ‰ to ca. +12 ‰ and ca. +16 ‰ respectively, which agrees with the observed  $\delta^{18}$ O values in sample #1598. The unchanged primary mineral association (like in sample #1734) experienced a complementary isotopic effect resulting in a decrease of the  $\delta^{18}$ O value.

The entire mineralogical-geochemical history of the inclusions can be envisaged as follows: Initially, within the lower mantle a calcite + CaSi-perovskite + volatile inclusion was encapsulated within diamond, forming a closed micro-system. During the ascent of the host diamond at ca. 15 GPa, CaSi-perovskite decomposed, with the formation of titanite (CaSi<sub>2</sub>O<sub>5</sub>), larnite (Ca<sub>2</sub>SiO<sub>4</sub>), walstromite (CaSiO<sub>3</sub>), and then, at ca. 4 GPa, wollastonite (CaSiO<sub>3</sub>) (Akaogi et al., 2004). During the final ascent at ca. 0.8 GPa, wollastonite, with the participation of the encapsulated fluid, dissociated into monticellite + cuspidine (Kushiro and Yoder 1964; Wirth et al. 2009). The oxygen isotope fractionation took place mostly during the latest transformations, at low temperatures (500–100  $^{\circ}$ C).

Some facts remain, however, unexplained. The suggested fractionation trend B (Fig. 4) with  $\pm 2$  % error lies outside of all known isotopic values for kimberlites, kimberlitic carbonates and primary igneous carbonatites (Fig. 4). In this case, the initial isotopic values for C and O in the lower mantle, as determined for the two analyzed samples, may be different to those of existing estimates for the upper mantle. Alternatively, the obtained data can be considered as the result of local fluctuations within a heterogeneous mantle (e.g., Rampone and Hofmann 2012). Further isotopic studies of lower-mantle minerals will help to explain better these unusual data.

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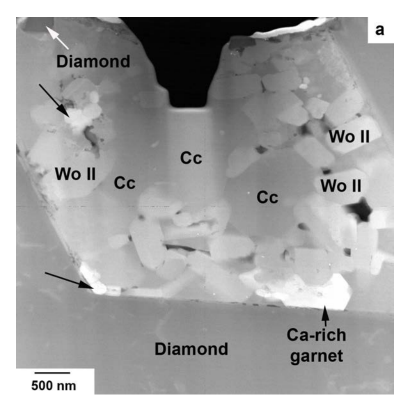
## **Table**

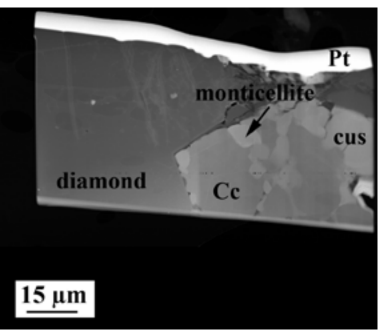
**Table 1.** Values of  $\delta^{18}$ O and  $\delta^{13}$ C in carbonate and silicate inclusions

Sample, mineral	$\delta^{18}$ O ± 1 $\sigma$ ,‰ SMOW	$\delta^{13}$ C ± 1 $\sigma$ ,‰ VPDB
Foil #1734		
Calcite	$-3.3 \pm 7.0$	$-2.8 \pm 6.0$
Wollastonite	-1.6 ± 8.2	
Foil #1598		
Calcite	+15.4 ± 4.3	+9.3 ± 6.1
Monticellite	+15.0 ± 6.5	

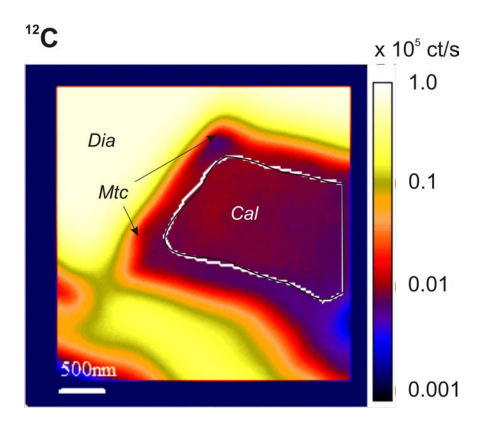
## **Captions to Figures**

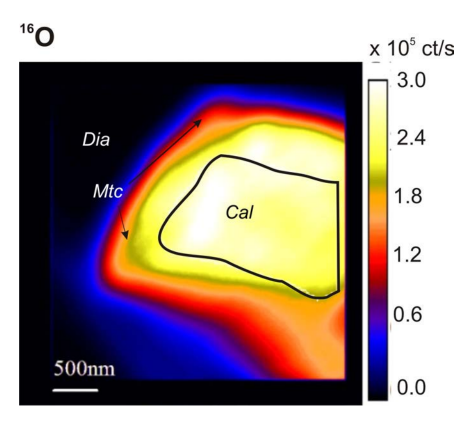
**Fig. 1.** HAADF TEM images of the studied associations. A – Calcite (Cc) + wollastonite (Wo II) association from inclusion in diamond #8/103, foil #1734. B – Calcite (Cc) + monticellite + wollastonite (wo) + cuspidine (Cus) association from inclusion in diamond #8/104, foil #1598. The bright spots in diamond indicated by white arrows are native iron (left) and iron + manganese (right).

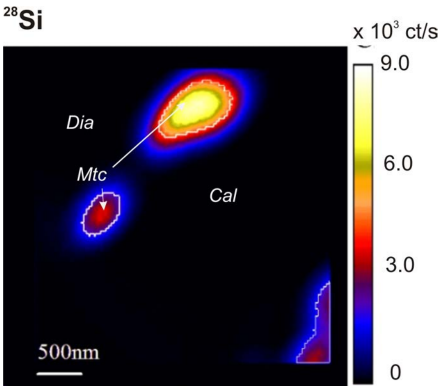




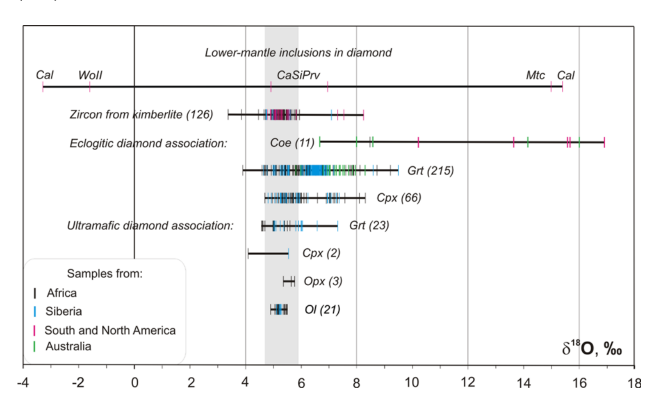
**Fig. 2** Ion images of the isotopic ratios of  $^{12}$ C (A),  $^{16}$ O (B) and  $^{28}$ Si (C) in sample #8/104 (foil #1598), rastered for  $4 \times 4 \ \mu m^2$ . White (black for  $^{18}$ O) lines outline the counted areas of the mineral grains. Dia – diamond, Cal – calcite, Mtc – monticellite. Color bars indicate yield of ions (intensity of response) in counts per one second (ct/s). The scales are set by the image processing software.







**Fig. 3** Values of  $\delta^{18}$ O for mantle minerals of diamond facies. Analyses of ultramafic association minerals after Mattey et al. (1994); Lowry et al. (1999); Spetsius et al. (2008); analyses of eclogitic association minerals after Deines et al. (1991); Jacob et al. (1994, 1998); Snyder et al. (1995); Lowry et al. (1999); Schulze et al. (2003, 2013); Viljoen et al. (2005); Spetsius et al. (2008); Ickert et al. (2013). Analyses of zircon from kimberlites after Valley et al. (1998) and Page et al. (2007). Number of analyses is shown in brackets at mineral names: OI – olivine, Opx – orthopyroxene, Cpx – clinopyroxene, Grt – garnet, Coe – coesite, Cal – calcite, WoII – wollastonite II, CaSiPrv – CaSi-perovskite, Mtc – monticellite. Data on lower-mantle CaSiperovskite from Harte et al. (1999). Shaded area is the "garnet mantle region" after Valley et al. (1998).



**Fig. 4** Isotopic composition of oxygen and carbon in lower-mantle minerals. Modified primary igneous carbonatite (PIC) box (red) after Srivastava et al. (2005); the field of carbonatites worldwide after Tappe et al. (2008); the upper-mantle  $\delta^{18}$ O area after Valley et al. (1998); the upper-mantle  $\delta^{13}$ C area after Cartigny (2005). Field of primary carbonates in kimberlites after Wilson et al. (2007). Explanations in text.

