The growth of lithospheric diamonds
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Natural diamonds contain mineral and fluid inclusions that record diamond growth conditions. Replicating the growth of inclusion-bearing diamonds in a laboratory is therefore a novel diagnostic tool to constrain the conditions of diamond formation in Earth’s lithosphere. By determining the carbon isotopic fractionation during diamond growth in fluids or melts, our laboratory experiments revealed that lithospheric monocrystalline and fibrous and coated diamonds grow similarly from redox reactions at isotopic equilibrium in water and carbonate-rich fluids or melts, and not from native carbon. These new results explain why most of the lithospheric diamonds are characterized by a common carbon isotopic fingerprint, inherited from their common parent fluids and not from the mantle assemblage.

**INTRODUCTION**

There is now a consensus that diamond growth in Earth’s upper mantle is related to carbon precipitation from mobile C-O-H-N-S-Cl fluids/melts, the detailed nature of which remains to be investigated (1–7). However, the associated carbon speciation (that is, redox state, carbonate melt or fluid, methane, elemental, or carbide) and origin (that is, mantle versus recycled and/or a mixture of both) are still debated (4–6). In particular, whether monocrystalline (gem-like) diamonds—and the so-called fibrous or coated diamonds—could form from the same parental medium is uncertain. Each of them would have markedly distinct genesis ages; the former are generally described as diamond xenocrysts in their host kimberlite, whereas the latter have short residence times (approximately on a scale of million years) in the mantle and are probably related to kimberlite magmatism (4, 5). Conversely, recent evidence has indicated a shared growth medium (7–9). Furthermore, fibrous diamonds record a wide variety of fluids that are saline, hydrous, and carbonatic in nature (1, 10). It remains unclear how these fluids are related. Some studies suggest that immiscibility processes (from a carbonatic fluid) would govern the chemical diversity (10). Others (11) suggest that a single original saline fluid would lead to two evolved fluids (carbonatic and hydrous silicic) depending on the chemical composition of the host rocks. These two opposing models are important to distinguish, as they propose distinct carbon source regions, namely, the asthenosphere and the subduction zones, respectively. C isotopes are central to the discussion and validation of these models, but how carbon behaves during diamond growth remains uncertain. For example, isotopic equilibration is generally assumed but has never been demonstrated (3). In natural systems, diamond generally lacks its original host rock, so addressing these issues is generally not possible. The study of synthetic inclusion-bearing diamonds, while difficult to perform, presents an alternative with which to circumvent this problem.

**RESULTS**

We produced syngenetic inclusion-bearing diamonds from fluids having different chemical compositions, real proxies for the peridotite-suite lithospheric diamonds and likely for the eclogite suite (9). Their carbon isotopic signature was measured at the micrometric scale using secondary ion mass spectrometry (NanoSIMS) (see Materials and Methods). The inclusion-bearing diamonds were grown on synthetic diamond seeds, from fluids under conditions representative of Earth’s lithosphere (that is, pressure, temperature, composition, and δ13C). These fluids are identical in chemical composition to the bulk composition of inclusions found in natural fibrous diamonds (1) and in some monocrystalline diamonds (11). Their composition includes up to 50 weight % water (pure, and saline up to 30 g/liter of NaCl), silicates, carbonates, and graphite (added as a second potential carbon source). Experiments were performed under pressures of 7 to 9 GPa and temperatures from 1200° to 1675°C (see the Supplementary Materials). Homogeneous diamond growth (that is, spontaneous nucleation) and heterogeneous growth (on seeds) were observed. The growth on seeds was significant (maximum oriented overgrowth from 4 to 10 μm in a few hours) and was associated with the trapping of multi-phased inclusions: minerals—including spontaneously grown diamonds (that is, diamond included in diamond: “diamond inclusions,” up to 3 μm in size at 1400°C; Fig. 1)—and fluids. The silicate assemblage trapped as inclusions in the newly grown diamond is also present in the solid matrix (that is, solid residue associated to the seeds after the experimental quench) associated to the diamonds. For sample H3913 (iron present), this assemblage is comparable to that observed in natural diamonds, such as olivine and CaMg carbonate (that is, dolomite) inclusions (table S1). These results support the hypothesis that mineral and fluid inclusions are syngentic with their host diamonds (4) (that is, formed simultaneously from the same melt or fluid). The syngenericity relationship between inclusions and diamonds is still highly debated today (12, 13), and the present experiments suggest synchronous trapping. Results show that monocrystalline and fibrous diamonds can be grown from the same parent fluids; they differ only in their habit because of varying supersaturation conditions (that is, growth conditions), as previously suggested (9, 14) and in agreement with the study of natural twinned diamond monocrystals having trapped their parent fluids at their interface (7). This is also supported by the presence of interfacial hydrous silicic fluid films between mineral inclusions and their host diamonds (15), interpreted as relics of the diamond parent fluids.

Diamonds with significant growth on seeds (Fig. 1) were cut with a focused ion beam (FIB) into 2- to 5-μm-thick slices containing inclusions, in order to expose the inclusion areas (Fig. 2). The carbon stable isotope composition of the starting materials was measured by conventional mass spectrometry, and that of the samples and starting seeds was measured by NanoSIMS. Table S1 shows the compositions of the slices as measured by NanoSIMS. The three investigated samples

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DISCUSSION

The $\delta^{13}C$ signature of a diamond depends on the carbon source and its speciation and is redox-sensitive (5). In nature, monocrystalline diamonds—including both fibrous and coated diamonds—fall within a narrow range of $\delta^{13}C$ of $-5 \pm 3\%$ (per mil), defining, together with other mantle products, mantle-derived carbon (5), not corresponding to either a surface carbonate [from $-3$ to $+4\%$ (5)] or organic matter [from $-45$ to $-15\%$ (5)]. Lithospheric diamonds from the eclogitic suite exhibit a significantly larger range, from approximately $-40$ to $3\%$ (5).

Some diamonds from the transition zone reach $\approx 25\%$ (5), which suggests recycled organic carbon at subduction zones as a potential carbon source for both eclogitic diamonds from the lithosphere and below (16). The bulk carbon isotope value for a given diamond may not always reflect those of the components that contributed to its formation. One single diamond can exhibit internal isotopic heterogeneities at the sub-millimeter scale (14, 17–19), attributable to either multistage growth from isotopically distinct or evolving sources (17, 18) or the progressive chemical and/or isotopic evolution of the fluid from which the diamond grew (14, 19).

The $\delta^{13}C$ compositions of some samples fall between the composition of the starting diamond seeds and the composition of the carbonate mixtures, which we attribute to an overlap during analysis. For samples H3913 and H3908 (7 GPa, 1400°C), respectively, the signature of a "new" diamond area ($\delta^{13}C = -17.9 \pm 4.4\%$) and of a diamond inclusion (that is, a diamond spontaneously grown in the fluid and subsequently trapped as an inclusion in diamond; $\delta^{13}C = -14.19 \pm 3.1\%$; see Fig. 2) was determined by quantitative isotope maps. These were compared to the bulk composition (that is, the average isotopic composition of the system) and calculated for all samples following the equation: $\delta^{13}C_{bulk} = x\delta^{13}C_{carbonate powder} + y\delta^{13}C_{graphite}$, where $x$ and $y$ are the weight fractions of both components loaded in the Pt capsules for each experiment, and where $x + y = 1$.

For sample HBD01, the experiment was performed at high temperature (1645°C). In this run, graphite participated in the diamond growth, resulting in a diamond $\delta^{13}C$ (~$-24.5$ to $-27.6\%$) very close to that of the graphitic source material (~$-26.35\%$). This result is in agreement with diamond growth experiments performed in the Fe-Ni-C system, with pure graphite as the carbon source (20), and is confirmed by the absence of residual graphite in the quenched products associated with the seeds. Although this experiment is not a relevant proxy for Earth’s mantle—being far from the geotherm—it illustrates the strong effect of temperature on the nature and behavior of the diamond parent fluid, making graphite available as a carbon source, resulting in a distinct isotopic...
composition. The lower-temperature diamond $\delta^{13}C$ compositions (1400°C) cannot originate from the graphite source ($-26.3 \pm 0.1\%$) by a direct dissolution of graphite in the fluid, followed by diamond growth, because the corresponding isotopic fractionation would be limited (5). For sample H3908, comparison of the C isotopic bulk signature ($-16.9\%$) with that of the diamond included in the diamonds ($-14.19 \pm 3.1\%$) shows that the inclinations could not have been generated from a fluid where all the available carbon (graphite and carbonates) would be dissolved [that is, a fluid with the composition of the bulk system (see Fig. 3 and table S3)]. Although we cannot rule out the possibility that a small amount of graphite could have been dissolved in the carbonate fluid or melt during the low-temperature experiments (1400°C), the presence of large graphite globules in the quenched products of samples H3913 and H3908 [up to 400 μm in diameter in 1 mm (diameter) × 1 to 2 mm (length) Pt capsules] implies that graphite is not significantly involved in diamond formation. The presence of dolomite (Ca,Mg carbonate) in the quenched products also demonstrates that carbonates were present in excess during the experiment, confirming that no carbon is needed from graphite to grow diamond. The absence of alkali carbonates in both the quenched matrix and as inclinations also suggests that these carbonates may act as major carbon sources.

The aqueous fluid and carbonate/silicate melt (8, 9) nature of the trapped inclusions also indicates the presence of two fluids in equilibrium with each other at high pressure and high temperature: a hydrous carbonate/silicate melt and an aqueous fluid, both enriched in carbon. If this were the case, the two fluids should be very close to miscibility at the pressure and temperature of the experiment (8), which means that they are close in composition and density, with all elements partitioning converging to unity (21). Therefore, as they should both exhibit the same $\delta^{13}C$ composition, we deduce that the isotopic composition of the parent fluid is that of the starting carbonated silicate powder (MELD, $-11.4\% \pm 0.1\%$).

Diamond growth is proposed to occur through redox reactions with COH fluids from either carbonate or CO$_2$ reduction (3, 14, 22) or from CH$_4$ oxidation (5, 23, 24) and through oxygen-conserving reactions in a water-rich fluid buffering the system and in which minor coexisting CH$_4$ or CO$_2$ reacts to form diamond by precipitation (25). Depending on the processes and sources involved during diamond growth, and because the fractionation between $^{12}C$ and $^{13}C$ is dependent on the processes and sources involved during diamond growth, and because the fractionation between $^{12}C$ and $^{13}C$ is dependent on the processes and sources involved during diamond growth, and because the fractionation between $^{12}C$ and $^{13}C$ is dependent on phase transformations processes and on carbon speciation in fluids, the carbon isotopes would be fractionated to different degrees. Core-to-rim variations of the $\delta^{13}C$ in diamonds are known indicators of redox processes: At 1000°C and under isotopic equilibrium, diamond is enriched in $^{13}C$ by approximately 2 to 3‰ compared to carbonate or CO$_2$ but is depleted in $^{13}C$ by approximately 1‰ when it precipitates from CH$_4$ (5). A reoxidation of Fe-Ni-C phases would involve a fractionation factor from 4.1 to 2.4‰ at 1450°C (20), whereas a precipitation process from dissolved carbon (no redox reaction) would cause a depletion of only a few tenths of a per mil (5). The diamond inclinations present in seeds from sample H3908 (Fig. 2) represent new uncontaminated diamond crystals. Therefore, we can use their isotopic composition to calculate the $\Delta_{\text{carbonatefluid-diamond}} = 2.8\%$, in agreement with diamond formation from fluid or melt: A calculation performed assuming isotopic equilibrium at 1400°C predicts that diamond should be depleted in $^{13}C$ by 2 to 4‰ compared to its carbonate or CO$_2$ source (5). This is also in agreement with diamond synthesis from pure sodium oxalate acting as the carbon source: $\Delta_{\text{carbonatefluid-diamond}} = 2.6\%$ at 7.5 GPa and 1400° to 1700°C (26). The low-temperature samples of the present study are relevant proxies for natural lithospheric diamonds, and their isotopic signatures show that all lithospheric diamonds, whether monocrystalline or fibrous, grow from oxidized carbon species dissolved in hydrous melt or fluids through redox mechanisms involving carbonate fluid or melt reduction (oxidized). For the present experiments, we propose the following two reacions (Equations 1 and 2):

$$12\text{Al}_2\text{O}_3 + 4\text{MgO} + K_2\text{CO}_3 + 16\text{SiO}_2 + 4\text{H}_2\text{O} + 2\text{CO}_2$$

$$= 4\text{K(Al, Mg)}_2(\text{OH})_2(\text{Si, Al})_O_{10}\text{phengite} + 4\text{C}_\text{Diamond} + 19\text{O}_2 \tag{1}$$

$$\text{Al}_2\text{O}_3 + 3\text{MgO} + 2(\text{Fe, Mg})\text{CO}_3 + 3\text{SiO}_2 + 2\text{H}_2\text{O} + 2\text{K}_2\text{CO}_3$$

$$+ 2\text{Na}_2\text{CO}_3 + 3\text{CaCO}_3 + 2\text{H}_2\text{O} + 2\text{CO}_2$$

$$= (\text{Fe, Mg})_2\text{Si}_2\text{O}_5\text{spinel} + \text{MgAl}_2\text{O}_4\text{spinel} + 10\text{C}_\text{Diamond}$$

$$+ 12\text{O}_2 + (\text{Ca, Mg})\text{CO}_3$$

$$+ 2(\text{Na, K})_2(\text{Ca, Mg})\text{Al}_2\text{Si}_2\text{O}_7\text{H}_2\text{O}_\text{Glass} \tag{2}$$

with H$_2$O and carbonates (possibly some aqueous CO$_2$ and/or carbonate groups CO$_3^{2-}$) being present in either the hydrous carbonatitic melt or the aqueous fluid (in chemical and isotopic equilibrium with each other).

Experiments performed under upper mantle conditions show that carbon isotopic fractionation is controlled by temperature, bulk composition, and the redox conditions (27). For experiment H3913 (iron present), olivine and diamond are produced; for this experiment, the starting materials were synthetic powders. We cannot calculate the
oxygen fugacity, but on the basis of the presence of olivine and dolomite, we can assume that $fO_2$ is close to FMQ-1 or FMQ-2 (fayalite + $O_2$ = magnetite + quartz), similar to EMOD (enstatite + magnetite = olivine + diamond). In experiments performed under comparable conditions (diamond seeds, silicate and carbonate powders, and water), but without graphite in the starting materials, no diamond growth is observed (that is, $fO_2$ conditions above EMOD). Therefore, the presence of graphite determines growth versus dissolution during the experiments, by buffering the redox of the system, acting as a $fO_2$ buffer and not as a C source. In the same fluid or melt, under less-reducing conditions (that is, with no graphite buffer present), diamond does not grow or is dissolved ($fO_2$ > 28). Under more-reducing conditions (that is, with a graphite buffer present), diamond grows ($fO_2$ < 22). This dependence on oxygen fugacity constrains the location of diamond formation in the mantle. Because the same primitive fluids and their immiscible products will lead to fibrous and monocrystalline diamond growth, they account for their common carbon isotopic composition.

A significant amount of carbon must be present in the upper mantle (29), possibly as fluids or melts located at the lithosphere-asthenosphere boundary (30, 31), before they are converted to diamonds when redox becomes favorable to growth.

MATERIALS AND METHODS
Sample preparation
FIB preparation was performed on a selection of diamonds: HBD01, H3908, and H3913 (fig. S1 and table S1). The diamonds were deposited on stubs covered with carbon tape. The selected diamonds were sliced open on one face and polished by the ion beam. Those that exhibited inclusions were sliced and ion beam-polished on the other face in order to obtain 2- to 5-μm-thick sections, including the trapped inclusions. The thick diamond sections were extracted and deposited with a FIB micromanipulator on silicon wafers and gold-coated for NanoSIMS measurements (fig. S2). The starting diamond seeds were embedded in indium and polished with the ion beam.

Analytical conditions and calibration for carbon isotope measurement with the NanoSIMS
Samples were coated with 10 nm of gold. The size (<10 μm) of the growth areas required a nanoscale analysis, which was only possible using the 150-nm NanoSIMS beam. Images were acquired using the NanoSIMS 50 (CAMECA) located at the Museum National d’Histoire Naturelle in Paris, France. Sample surface was sputtered by a 1.5-μA Cs beam to obtain images of various sizes from 5 μm × 5 μm to 30 μm × 30 μm and divided into 256 pixels × 256 pixels. Raster speed was regulated at 1 ms per pixel, with an approximate spatial resolution of 150 nm. Secondary images of $^{12}C$, $^{13}C$, $^{16}O$, and $^{28}Si$ were simultaneously collected. Before each acquisition, the sample surface was presputtered over a 30 μm × 30 μm region using a 100-μA Cs+ beam for 10 min to remove gold coating and surface contamination and to reach sputtering steady state. The images were processed using the L’IMAGE software (L. Nittler, Carnegie Institution). $^{13}C$ and $^{15}C$ were used to generate $\delta^{13}C$ isotopic compositions, relative to the PDB (Pee Dee Belemnite) standard. Regions of interest were manually drawn to isolate new diamond growth areas and to determine their isotopic composition. The calibration was performed against broken pieces of natural Ia and a synthetic Ia diamond used for diamond anvil cells, whose compositions were determined by gas source mass spectrometry at IPGP (Institut de Physique du Globe de Paris) at −3.6 ± 0.1‰ and −20.9 ± 0.1‰. These carbon isotope compositions were determined by combustion of each diamond, followed by analysis of the resulting CO$_2$ gas using a Thermo Fisher Delta + XP dual-inlet gas source mass spectrometer. Diamonds were individually wrapped in Pt foil and combusted under vacuum at 1100°C in pure oxygen. Extracted CO$_2$ gas was purified and gas pressure–quantified for yield evaluation under cryogenic conditions before collection and analysis. The carbon isotopic values obtained for the diamonds are reported as $\delta^{13}C = \left(\frac{^{13}C}{^{12}C}_{\text{sample}}\right)_{V-PDB} - 1) \times 1000$, with an uncertainty of <0.06‰ (2σ).

The calibration methodology is presented in the study of Pinti et al. (32).

The starting materials (powders) used for the experiments were similarly characterized for their $\delta^{13}C$ using gas source mass spectrometry at GEOTOP, Montréal, using three reference standards at −42.16, −28.75, and −11.85‰, respectively, versus VPDB (Vienna Pee Dee Belemnite) with an uncertainty of 0.1‰.

SUPPLEMENTARY MATERIALS
Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/4/6/eaat1602/DC1

table S1. Isotopic compositions of the starting materials, standards, and samples.
table S2. Description of the studied samples.
table S3. Mass balance calculations of isotopic compositions of the bulk composition for the starting materials and the high-pressure and high-temperature carbonate fluid.
fig. S1. SEM images of the FIB preparation, cut on one side of the seed and on the other side to obtain the final slices.
fig. S2. Details of the samples analyzed with the NanoSIMS.
fig. S3. NanoSIMS maps of $^{12}C$ for each investigated sample showing the locations of the areas of interest (red squares) corresponding to the location of the measurements reported in table S1 and Fig. 1.

Reference (33)

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