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CaSiO₃-perovskite in diamond confirms the recycling of oceanic crust into the lower mantle

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Laboratory experiments and seismology have created a clear picture of the major minerals expected to comprise the deeper parts of Earth's mantle. Finds of some of these phases in superdeep diamonds have confirmed part of this picture [1-5]. A striking exception is the high-pressure perovskite structured polymorph of CaSiO₃. This mineral – inferred to be the fourth most abundant in Earth - has never been found in nature. As the major host for Ca and heat-producing elements K, U and Th in the transition zone and lower mantle, it is critical to establish its presence. Here we document the first discovery of the perovskite-structured polymorph of CaSiO₃ in nature, included within a diamond from the Cullinan kimberlite, South Africa. The mineral is intergrown with ~ 6% CaTiO₃. The Ti-rich nature of this association indicates a bulk composition consistent with derivation from basaltic oceanic crust subducted to pressures equivalent to depths of the uppermost lower mantle. The relatively heavy carbon isotopic composition of the surrounding diamond, δ^{13} C = -2.3 \pm 0.5%, together with the pristine high-pressure CaSiO₃ structure, provides evidence of the recycling of oceanic crust and surficial carbon to lower mantle depths.

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A key goal of solid Earth geosciences is to establish the mineralogy of Earth's mantle throughout its depth as it acts a primary control on mantle dynamics and chemistry. Diamonds are unique windows in this regard, providing access to the deepest intact material from Earth's interior through included minerals. Over three decades, a growing number of studies have used a class of diamonds known as "super-deep" diamonds to study mantle processes in the deep sublithospheric mantle, the transition zone and the lower mantle1-6. Pioneering studies1-3 suggested that some of the assemblages included within super-deep diamonds represented samples of the lower mantle and transition zone, variably retrogressed to lower pressures. Later studies cautioned that some of these assemblages and minerals might originate at shallower depths^{7,8}, though still beneath the lithosphere.

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The most common minerals found within super-deep diamonds are ferropericlase [(Mg,Fe)O] and CaSiO₃^{1-3,9}. Ferropericlase is stable at most pressure and temperature conditions in the mantle and hence, when found as a single inclusion within diamond, this mineral cannot be considered an unambiguous indicator of a super-deep origin⁷.

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The CaSiO₃ phase found within super-deep diamonds typically has the crystal structure of walstromite (BaCa₂Si₃O₉)^{1,6,8,9}. Perovskite-structured CaSiO₃ (Ca-Pv) is considered one of the most important components in the Earth's lower mantle, comprising approximately 7% of a peridotitic mantle composition and ~23% of the volume of a subducted mid-ocean ridge basalt slab⁹⁻¹¹. As such, it is likely to be the fourth most abundant terrestrial mineral. Within a peridotitic lower mantle, Ca-Pv is the dominant sink for Ca and for incompatible elements, including the key heatproducing elements K, U and Th¹². However, Ca-Pv has so far never been found in nature and even high-pressure laboratory experiments have failed to quench it as a metastable phase at the Earth's surface. While initial studies of super-deep diamonds made a clear case for the presence of Ca-Pv¹⁻ 4, their structure was either undetermined or documented to be the lower-pressure polymorph -CaSiO₃-walstromite - and interpreted as a back-transformation of perovskite-structured CaSiO₃. The CaSiO₃-perovskite to CaSiO₃-walstromite phase transformation would require a volume change¹³ of about 28% that is impossible for diamond to accommodate due to its extremely high bulk modulus¹⁴. The absence of healed fractures in the diamond host studied by ¹³ implies that CaSiO₃-walstromite, in that case, is unlikely to represent inverted CaSiO₃-perovskite. Plastic deformation of the diamond lattice could accommodate some of the volume change necessary for

the inclusion's phase transformation. Although plastic deformation is well documented in super-deep diamonds¹⁵, it has never been quantified and the amount of deformation would have to be substantial. Thus, though some super-deep diamonds with documented phase assemblages that include ferropericlase, enstatite (inverted bridgmanite) and/or CaSiO₃-walstromite are likely to have originated from lower mantle depths^{1-4,9}, there remains some ambiguity hence finding an unretrogressed silicate perovskite would provide very powerful confirmation of lower mantle sampling by some super-deep diamonds.

Here we investigated an inclusion within a diamond from the Cullinan Mine, Gauteng Province, Republic of South Africa. The Cullinan kimberlite is Group I in character, i.e., its chemistry and Sr-Nd-Hf isotope signatures are thought to reflect a melt source from beneath the lithospheric mantle, within the Earth's convecting mantle¹⁶. Cullinan mine is renowned for producing exceptionally large diamonds such as the Cullinan diamond^{6,17}, most of which have been suggested to be super-deep diamonds⁶.

The 31×26×10 µm³ CaSiO₃ inclusion in diamond, exposed by polishing, was revealed by X-ray diffraction, Raman spectroscopy and electron backscatter diffraction to have a perovskite structure. To our knowledge, this represents the first finding of a non-reverted silicate perovskite-structured CaSiO₃ phase found in nature and the first, including those synthesized in the laboratory, to be preserved with its high-pressure structure at the surface of the Earth.

Cathodoluminescence imaging of the host diamond surrounding the Ca-Pv inclusion (Fig. 1) reveals multiple growth zones and a complex internal structure, typical of super-deep diamonds^{4,18}. FTIR spectroscopy (Extended Data Figure 1) of the diamond host indicates a nitrogen content of 34 ppm, with 97% in the B-aggregated form, i.e., the diamond host is type IaB. The low nitrogen content and very high level of B aggregation are typical characteristics of other super-deep diamonds^{4,19}

94 indicating prolonged residence at the high temperatures expected at transition zone and lower 95 mantle depths.

The chemical composition of the CaPv inclusion, determined by electron microprobe, yields a nearly pure CaSiO₃ composition (Ca_{0.98}Si_{0.98}O₃), with minor impurities of Ti, Al, Fe, Mg totaling 0.04 atoms per formula unit (Extended Data, Table 1).

Backscatter electron imaging and EDS element maps (Fig. 2) show that the Ca-Pv crystal includes 14 irregular areas of CaTiO₃-perovskite between 1 and 7-8 µm with a stoichiometry of ~ Ca(Ti_{0.92}Si_{0.07}Al_{0.02})O₃. The size and abundance of the CaTiO₃ intergrowths within this Ca-Pv is very similar to the texture and abundance of inclusions reported in CaSiO₃-walstromite phases reported from Juina super-deep diamonds²⁰. The 2D exposed surface of our Ca-Pv inclusion makes accurate estimation of its bulk composition difficult, but image analysis indicates that the host crystal, in bulk may have up to 6% by volume CaTiO₃. CaTiO₃-perovskite is a common mineral in nature and is stable well into the lower mantle²¹. In contrast, CaPv retaining its perovskite structure has no experimentally synthesised analogues at room temperature and pressure, unless significant CaTiO₃ is dissolved within its structure (~34 mol %²¹), far more than the CaTiO₃ component observed here. However, our discovery of natural CaPv, trapped in a diamond, with < 2 mol % CaTiO₃ component in the CaSiO₃-rich portion of the inclusion, indicates that unlike experiments, there must be natural P-T-t pathways that are capable of the metastable preservation of this phase.

X-ray diffraction data show the CaSiO₃ inclusion to have the perovskite structure. The small size of the inclusion (thickness estimated by confocal Raman spectroscopy to be $\leq 10 \mu m$) and its entrapment within the diamond host resulted in only a limited number of measured diffraction reflections (n=91), of which only 9 were unique (Extended Data Table 2). The full 91 reflections were used to refine the Ca-Pv unit-cell parameters to yield:

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However, alternative unit-cell refinements using other numerical approaches could provide significantly different unit-cell parameters that may differ by greater than 1%, due to the relatively poor accuracy and precision of the d-spacings measured here. These larger than normal uncertainties are typical when studying minerals of this size and arise not just from the limited number of reflections but also due to the measurements being performed using an area detector, which provides lower precision in d-spacing determination than a point detector. Such relatively large uncertainty on the cell parameters makes any comparison with the unit cell volume of CaTiO₃ -perovskite unreliable, even though we can define the two structures as being very similar. Ewald projections along the three crystallographic axes (Fig. 3a) indicate an orthorhombic unit-cell. The unit-cell together with the chemical composition define the mineral to be perovskite-structured CaSiO₃. Recent numerical simulations on "host-inclusion" systems²², indicate that an inclusion partly exposed to atmospheric pressure only loses a portion of its residual pressure, as a function of the elastic properties of both the host mineral and the inclusion. The Ca-Pv inclusion studied here is partly exposed at the diamond surface but with 2/3 of its volume still buried in the diamond host. Thus, any measurements on this grain would be affected by some residual "pressure" still acting on the inclusion, which in turn affects the X-ray diffraction data and Raman spectra.

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Raman spectra (Fig. 3b) for the inclusion show that the CaTiO₃ perovskite spectrum is in excellent agreement with CaTiO₃-perovskite Raman data from the RRUFF database²³ (Extended Data Fig. 2). The CaSiO₃ and CaTiO₃ spectra are similar. Small differences are evident due to the presence of two Raman peaks for the CaSiO₃ spectrum, which could belong to the lower-pressure CaSiO₃ polymorph wollastonite-2M²³. This wollastonite polymorph is not stable at pressures higher than 3 GPa along a mantle geotherm ²⁴, well below the diamond stability field. Hence its presence seems likely due to minor partial inversion of the Ca-Pv phase owing to the polishing of the sample to expose the inclusion, as reported in previous work²⁵.

The EBSD measurements collected on several areas of the grain provide no evidence that amorphous portions are present and confirm that the CaSiO₃ zones have perovskite-structure (Fig. 4). A typical EBSD pattern collected on the CaSiO₃ area (red circle), shown as the relative non-indexed EBSD pattern (Fig. 4) is complex and could not be indexed by a single phase. Indexing the pattern by using a combination of reference EBSD patterns for CaTiO₃-perovskite (Fig. 4c) and wollastonite-2M (Fig. 4d), accounts for the observed pattern, confirming the conclusions from X-ray and Raman work, that CaSiO₃ is present in this diamond with a perovskite-type structure.

We suggest that the natural Ca-Pv found trapped within our super-deep diamond originated as a result of unmixing, of the high-pressure solid solution $Ca(Ti,Si)O_3$. If the two phases exsolved from a homogenous bulk composition, this phase would contain $\sim 3.9\%$ TiO₂.

Our estimate of the stoichiometry of the original phase composition is:

 $(Ca_{0.98}Mg_{0.01}Fe_{0.01})(Si_{0.93}Ti_{0.06}Al_{0.01})O_{3}$

This composition is consistent with those of $CaSiO_3$ crystallised in experiments from a MORB-like bulk composition at $\sim 24~GPa^{26}$ and is similar to the $CaSiO_3$ - walstromite/CaTiO $_3$ intergrowths found within Juina super-deep diamonds that were proposed to represent basalt-like compositions subducted to lower mantle depths and later retrogressed during ascent to Earth's surface²⁰. The preservation of the high-pressure perovskite structure in the case of the Cullinan inclusion proves the derivation of such compositions from lower mantle depths.

The indicated subducted basaltic protolith of the Cullinan Ca-Pv inclusion suggests that we might expect to observe some evidence of a crustal parentage in the C isotopic composition of the host diamond. Its carbon isotopic composition is variable (Fig. 1; Extended Data: Table 3) with δ^{13} C values ranging from -2.3‰ to -4.6‰. The core region of the diamond, defined by CL-imaging (Fig. 1), contains the Ca-Pv inclusion and an average δ^{13} C value of -2.3 ±0.5‰, significantly lower than

the typical upper mantle value of -5.5%²⁷. In contrast, the outer rim region of the diamond has a composition (Mean δ^{13} C -4.1 \pm 0.5‰) that is closer to the normal mantle value. Crustal carbon reservoirs have C isotopic compositions that are both heavier and lighter than the typical upper mantle value. While isotopically lighter carbon (below -25%) has been found in super-deep diamonds from Juina, proposed to be derived from subducted basalt protoliths^{20,28}, heavier carbon isotopic compositions, such as those measured in the core of our Cullinan diamond, have also been reported in super-deep diamonds from both Sao Luis/Juina (Brazil) and from Kankan (Guinea)¹⁸⁻²⁰. If the δ^{13} C value of -2.3% is compared with the median value (-4.91 %) of 1473 published analyses of lithospheric diamonds containing peridotitic inclusions – a group of diamonds usually accepted to have minimal subducted influence²⁷ - it can be defined as an outlier, beyond 3 times the median absolute deviation. Such anomalously heavy C isotopic compositions are thought to reflect a greater influence of subducted carbonate in the fluid that formed these super-deep diamonds 18,19. The C isotope compositions of the rim of the Cullinan diamond (Fig. 1) may represent an overgrowth that grew under upper mantle conditions, or from a distinct source of carbon in the lower mantle. Regardless, the isotopically heavy δ^{13} C values of the portion of the diamond containing the Ca-Pv inclusion lends support to its origin from a subducted basaltic protolith.

Our discovery of perovskite-structured CaSiO₃ in a super-deep diamond firmly establishes this phase as a component in Earth's deep mantle, confirming previous suggestions that lower pressure CaSiO₃ polymorphs included in these diamonds may represent retrogressed Ca-Pv^{1-4,9}. The estimated original bulk composition of the Cullinan Ca-Pv inclusion is consistent with compositions stable in subducted oceanic basalt protoliths at ~ 24 GPa, in the uppermost lower mantle²⁶. Our finding thus confirms the expectation from calculation¹⁰ and high-pressure experiments^{21,25}, that Ca-Pv is the chief Ca-bearing phase in the lower mantle in both basic and ultrabasic compositions, reaching up to 23 vol% in MORB-like compositions²⁶. The combined bulk composition of the Ca-Pv phase found here provides overwhelming evidence of the return of recycled oceanic crust into

- Earth's lower mantle²⁰ while the relatively heavy C isotopic composition of the diamond in contact
- with the inclusion indicates the subduction of crustal carbon to lower mantle depths.

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AUTHOR CONTRIBUTIONS

FN conceived the study, wrote the initial manuscript and performed X-ray diffraction and micro-Raman measurements. NK found the mineral, made original mineral identifications on a confocal Raman spectrometer (EOAS, UBC), performed microprobe and CL measurements, prepared samples for SIMS measurements and assisted with the manuscript preparation. MK supervised the study of the Cullinan diamond collection acquired through JG, AEM and JD and assisted with manuscript preparation. DGP made the geochemical interpretations and led the manuscript revisions. MGP assisted with the manuscript preparation and crystallographic interpretations. NR, MGP and MA assisted with the X-ray data interpretation. LP collected and interpreted the EBSD data. JG, AEM and JD designed the sampling program.

227 COMPETING INTERESTS

The authors declare no competing financial interests.

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Perspective. Annu. Rev. Earth Planet. Sci. 42, 699–732 (2014). 297 298 28. Burnham, A.D. et al. Stable isotope evidence for crustal recycling as recorded by superdeep diamonds. Earth Planet. Sci. Lett. 432, 374-380 (2015). 299 300 301 302 303 Figure captions 304 305 Figure 1. Cathodoluminescence image and carbon isotopic composition of the diamond containing the 306 307 CaSiO₃ (Ca-Pv) inclusion. Inclusion of Ca-Pv shown in yellow. Spots at five different locations give carbon isotopic compositions of the diamond host, in δ^{13} C notation, where 308 δ^{13} C = $[(^{13}C/^{12}C)_{\text{sample}}/(^{13}C/^{12}C)_{\text{PDB}} - 1]x1000$ in which PDB is the Pee Dee Belemenite reference material. 309 Figure 2. Back-scattered electron image of the CaPv inclusion still included in host diamond and 310 energy-dispersive X-ray spectroscopy elemental maps. Back-scattered electron image (a) of CaPy inclusion 311 (dark-grey) surrounded by the diamond host (black) showing inclusions of CaTiO₃ perovskite (light gray). 312 Images b, c and d are energy-dispersive X-ray spectroscopy elemental maps of Ca, Ti and Si, respectively, 313 with intensity of colour (black within grain outline though to saturation in specific colour) proportional to 314 element concentration. 315 316 Figure 3. Ewald projections of X-ray diffraction data and Raman spectroscopic results. The Ewald 317 projections (a) were made along three different orientations for CaSiO₃-perovskite. The projections were obtained using Crysalis software (Rigaku-Oxford Diffraction). Baseline corrected Raman spectra in (b) 318 compare spectra from the CaPv inclusion with that of CaTiO₃ - perovskite found as an intergrowth (Fig. 2a). 319 320 Raman spot size was $\sim 1.1 \, \mu m$, spectral resolution 3 cm⁻¹. Figure 4. Electron backscatter diffraction (EBSD) images of the Ca-Pv inclusion in diamond. EBSD image 321 322 collected on the red circle (a) relative to the CaSiO₃ area of our inclusion. In (b) the non-indexed EBSD pattern is shown, whereas in (c) and (d) the pattern was indexed with the CaTiO₃ and wollastonite-2M 323 324 reference patterns. 325 **On-line Methods** 326 327 328 Micro-Raman spectroscopy 329 The CaSiO₃-perovskite sample was analysed by a InVia Renishaw micro-Raman spectrometer installed at 330 the Department of Chemical Sciences, University of Padova. Spectra were baseline corrected. A 632.8 nm 331 excitation laser was used at a power of 7 mW. The Raman spectrum of the CaSiO₃-perovskite crystal was 332 collected for 40 seconds using a 50× objective with a spatial resolution of 1.1 µm and a spectral resolution

estimated to be ~ 3 cm⁻¹. The most intense Raman peaks observed for the CaSiO₃-perovskite inclusion are, in

order of decreasing intensity (in cm⁻¹): 774, 247, 470, 337, 181 and 226.

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A direct comparison between the Raman spectrum of natural CaSiO₃-perovskite with that of CaTiO₃ inclusions and those reported by the RRUFF Raman database²³ indicate that the two spectra are very similar. A small but important difference is due to the presence of limited traces of wollastonite 2M on the natural CaSiO₃-perovskite Raman spectrum (see peaks at 971 and 637 cm⁻¹), which, as expected, are not evident for CaTiO₃ perovskite inclusions.

In terms of the Raman peak assignment, it must be noted that based on the results of [29], the broad Raman bands in the 650-850 cm⁻¹ region are due to second order Raman scattering and that only the sharp peaks in the 200-500 cm⁻¹ region are first-order Raman bands. For our purpose, however, the entire Raman spectrum of natural CaSiO₃-perovskite is considered, regardless of the first or second order scattering, for a direct comparison mainly with CaTiO₃ perovskite and wollastonite.

The strong similarity in the Raman spectra between $CaTiO_3$ and $CaSiO_3$ perovskites in Fig. 3b could be used to invoke the possibility that the spectra are dominated by a larger underlying, unexposed portion of the $CaTiO_3$ phase and also permit the possibility that the $CaSiO_3$ regions of the inclusion are amorphous, as we would expect a general Raman shift for the $CaSiO_3$ bands toward higher wavenumbers. This possibility can be discounted for a number of reasons. Firstly, the partially exposed inclusion is under some stress and this will affect the Raman band shift depending on the elastic properties of the two perovskites. More importantly, the Raman spectra of such a large amorphous area of $CaSiO_3$ would be totally distinct from that measured here (Fig. 3b), and would be, in such a scenario, characterized by the presence of three very intense Raman bands at ~ 370 , 640 and 970 cm⁻¹ (depending on the P and T conditions^{30,31}). These Raman bands are absent in the spectrum of the $CaSiO_3$ portion of the perovskite-structured inclusion in diamond. Also, the spot size and confocal nature of the Raman measurements made here are too small to be significantly influenced by the spatially associated $CaTiO_3$ intergrowth.

Cathodoluminescence

The cathodoluminescence SEM image (CL) shown in Figure 1 of the main text was obtained using a Philips XL 30 scanning electron microscope with a CL attachment consisting of a Hamamatsu R376 photomultiplier

tube (EOAS UBC, Vancouver, Canada). The accelerating voltage was 20 keV and the electron beam current was $100~\mu A$.

Infrared spectroscopy

Infrared spectra were collected for the diamond hosting natural CaSiO₃-perovskite on a Nicolet 6700 Fourier transform infrared spectrometer (Vancouver, Canada). The absorbance spectra for a sample were measured at maximum light transmission. Background spectra were collected for 120 s prior to the analysis and were subtracted from each measured absorbance spectra. Count times for spectra were 40 seconds at a spectral resolution of 0.5 cm⁻¹. The nitrogen concentration and aggregation were determined by the procedure described in [32] using the spreadsheet ("FTIR analyser 3d") provided by John Chapman (Rio Tinto Diamonds Ltd.). Preliminary processing and baseline determination were made using EssentialFTIR® software. The analytical and processing error is ±10% (1 sigma, relative error). The FTIR spectrum of the diamond from Cullinan studied here is shown in Extended Data Fig. 1.

Electron Microprobe analysis

Quantitative chemical analyses were undertaken on a fully automated CAMECA SX-50 electron microprobe (University of British Columbia, Department of Earth, Ocean and Atmospheric Sciences), operating in the wavelength-dispersion mode with the following operating conditions: excitation voltage, 15 kV; beam current, 20 nA; peak count time, 20 s; background count-time, 10 s; actual spot diameter, 5 μ m. Data reduction was done using the 'PAP' $\phi(\rho Z)$ method³². Detection limits for most oxides were below 0.08 wt.%, detection limits for Cr₂O₃, MnO₂, and NiO were less than 0.12 wt.%. Due to the crystal size of natural CaSiO₃-perovskite and the presence of inclusions of CaTiO₃ perovskite, we were able to perform only three reliable analyses; the results are reported in EXTENDED DATA Table 1. Na and K were not analysed.

Scanning Electron Microscopy and Energy Dispersive X-ray Spectroscopy

Our newly discovered natural CaSiO₃-perovskite was studied by SEM-EDS in order to investigate the distribution of Ca, Si and Ti over the grain. We used a CamScan MX3000 electron microscope equipped with a LaB6 source, four-quadrant solid state back-scattered electron (BSE) detector and an EDAX EDS

system for micro-analysis installed at the Department of Geosciences, University of Padova. The analytical conditions were: accelerating voltage, 20 kV; filament emission, 13 nA; working distance, 27 mm. The BSE image and the relative EDS Ca-Si-Ti map are shown in Figure 2.

Single-crystal micro-X-ray diffraction

Single-crystal X-ray diffraction measurements were performed using a Rigaku-Oxford Diffraction Supernova goniometer installed at the Department of Geosciences, University of Padova, equipped with a Dectris Pilatus 200 K area detector and a Mova X-ray microsource (MoK α -radiation) operating at 50 kV and 0.8 mA. The sample to detector distance was 68 mm. Data reduction was performed using the CrysAlis software (Rigaku Oxford Diffraction). Diffraction data are reported in Extended Data Table 2 and are compared to those of a reference CaTiO₃ measured on a single crystal³⁴ having the following unit-cell parameters: a = 5.388(1) Å, b = 5.447(1) Å, c = 7.654(1) Å, V = 224.63(1) Å3.

Carbon isotope analyses

Carbon isotope compositions (δ^{13} C), reported in EXTENDED DATA Table 3 were determined using a Cameca IMS 7f-GEO secondary ion mass spectrometer (Caltech, USA). The polished diamond investigated was pressed into an indium mount with a 1" diameter aluminium holder. Natural reference diamonds with reference values of δ^{13} C between -13.6% ($2\sigma = 0.3$) and -2.5-2.6% ($2\sigma = 0.3$) were used to determine the instrumental mass fractionation and drift before and after sample analyses. Diamonds were coated with gold (20 nm). Analyses were conducted using 133Cs^+ at 10 keV impact energy and a beam current of ~4 nA. The 15 μ m diameter 133Cs^+ primary-ion beam was used for pre-sputtering. During analysis, the ion beam diameter was reduced to 5 μ m. Secondary ions of 12C and 13C were extracted at -9 keV. No e-gun charge compensation was required. The secondary ion energy bandwidth was 90 eV. 13C-/12C- ratios were measured using dual Faraday cups (FC1 for 12C- and FC2 for 13C-). The mass resolving power (R = M/ Δ M) was 2900. The 12 C as well as 13 C ions were counted for 1 second, in each cycle (30 cycles in total). Total time spent on each spot was 8 minutes. The standard deviation of the analyses is estimated at about 0.4 to 0.5 % at the two-sigma (95% uncertainty) level.

420 Electron Backscatter Diffraction

- 421 EBSD analyses were performed at CNR-ICMATE (Padova, Italy) using a Quanta 200F FEG-ESEM (FEI
- Company) operating in high vacuum mode at an accelerating voltage of 30 kV, emission 174 μA, spot 4.5,
- without any conductive coating. EBSD patterns were collected at 10 mm WD and 75° specimen tilt, using an
- 424 EDAX Digiview EBSD system. The instrument is controlled by the OIMTM 5.31 software, which contains a
- 425 large EBSD pattern database.

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Statistical analysis of diamond carbon isotope composition

- 428 We used a compilation of 1473 C-isotope analyses of diamonds containing inclusions of lithospheric
- 429 peridotite paragenesis, from the dataset used by Huber (1981) [35]. We calculated the median absolute
- 430 deviation for this dataset using a bespoke Excel spread sheet, using formulae given in [36], using a b factor
- of 1.4826 and a very conservative threshold factor of 3 ³⁶.

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References – On-line Methods

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Data Availability Statement All relevant data are included in the Extended Data section of this manuscript (Extended Data Tables 1-3, forming the basis of Figs 1-4, except original spectral data and electron microprobe data which are available from the first author. **Extended Data Table and Figure Legends EXTENDED DATA Table 1.** Chemical analyses for CaSiO₃-perovskite investigated in this work, including cation proportions (calculated on the basis of 3 oxygens) for the average value (Ave). The data were averaged for three spot analyses due to the extremely limited crystal size. **EXTENDED DATA Table 2.** List of d-spacings with their relative intensities (I) with the most intense peak at 100 and hkl indexes for CaSiO₃-perovskite obtained by single-crystal X-ray micro diffraction compared with the *d*-spacings from a reference CaTiO₃ [34]. **EXTENDED DATA Table 3**. Carbon isotopic composition (δ^{13} C, in ‰) and relative uncertainty for the host diamond enclosing the CaSiO₃-perovskite inclusions. The locations from 1 to 5 refer to positions annotated on Figure 1 of the main text. **EXTENDED DATA Figure 1.** FTIR absorption spectrum, baseline corrected, of the diamond containing the CaSiO₃-perovskite inclusion. EXTENDED DATA Figure 2. Comparison between the Raman spectra of CaTiO₃ studied in this work (blue) and CaTiO₃ reported in the RRUFF database²³ (card number R050456). Footnote of the EXTENDED DATA Table 1 Cations calculated from data in column 2. Na and K were not analysed. Ave = Average. Stand. Dev. is the standard deviation of the 3 analyses used to construct the average in column 2. < D.L. denotes less than the detection limits listed in METHODS.







