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A primary natrocarbonatitic association in the Deep Earth

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Abstract

In addition to ultramafic and mafic associations, a primary natrocarbonatitic association occurs in the lower mantle. To date, it was identified as inclusions in diamonds from the Juina area, Mato Grosso State, Brazil. It comprises almost 50 mineral species: carbonates, halides, fluorides, phosphates, sulfates, oxides, silicates, sulfides and native elements. In addition, volatiles are present in this association. Among oxides, coexisting periclase and wüstite were identified, pointing to the formation of the natrocarbonatitic association at a depth greater than 2,000 km. Some iron-rich (Mg,Fe)O inclusions in diamond are attributed to the lowermost mantle. The initial lower-mantle carbonatitic melt formed as a result of low-fraction partial melting of carbon-containing lower-mantle material, rich in P, F, Cl and other volatile elements, at the core-mantle boundary. During ascent to the surface, the initial carbonatitic melt dissociated into two immiscible parts, a carbonate-silicate and a chloride-carbonate melt. The latter melt is parental to the natrocarbonatitic lower-mantle association. Diamonds with carbonatitic inclusions were formed in carbonatitic melts or high-density fluids.

Keywords: carbonatite, Deep Earth, lower mantle, carbonates, halides, fluorides, phosphates.

Introduction

According to current models, the Deep Earth has an ultramafic composition. It comprises an association of oxides: ferropericlase + bridgmanite (former MgSi-perovskite; Tschauner et al., 2014) + CaSi-perovskites (along with some other minerals), which replace, at depths of *ca.* 660 km, the silicate association of ringwoodite and majoritic garnet (e.g., Stixrude and Lithgow-Bertelloni 2007). In addition to these mineral phases occur minor amounts of carbonates, evidenced by singular grains of calcite and magnesite that have been reported from both upper-mantle (Sobolev et al. 1997; Wang et al. 1996) and lower-mantle (Brenker et al. 2007) associations. Multiple finds of saline or hydrous-saline fluid inclusions in fibrous diamond (e.g., Smith 2014 and references therein) further imply the presence of carbonatitic media during the formation of diamond in the mantle. Recently, a series of carbonatitic minerals was identified as inclusions in lower-mantle diamond from Juina in Mato Grosso State, Brazil (Wirth et al. 2009; Kaminsky et al. 2009, 2013, 2015a) confirming the presence of carbonatitic mineral associations in the Deep Earth.

On the other hand, experiments under high pressure indicate a significant role of carbonatitic media in the formation of diamond. For example, ‘super-deep’ diamonds have been experimentally crystallized in melts of the

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lower mantle, diamond parental carbonate - magnesio-wüstite – bridgmanite – carbon system at pressures of up to 60 GPa (Litvin et al. 2014).

In this work we summarize the existing knowledge on the carbonatitic mineral association in the lower mantle, along with presenting some newly obtained data, and an analysis of the conditions and the source(s) of the carbonatitic material in the Deep Earth.

Lower-mantle carbonatitic association in diamonds from the Juina area, Brazil

To date, almost 50 mineral species of the lower mantle carbonatitic association have been identified as inclusions in diamonds from the Juina, Mato Grosso State, Brazil (Table 1), which preserved them from various dissociation reactions that likely would have occurred during ascent to the surface. Some of the carbonatitic association species occur in union with lower-mantle minerals of the ultramafic association, such as CaSi-perovskite (Brenker et al. 2007) and ferropericlase (Kaminsky et al. 2009). This close association is strong evidence of their formation within the Deep Earth.

The following mineral classes can be distinguished among these minerals.

1. Carbonates.
2. Halides, fluorides, phosphates and sulfates.
3. Oxides.
4. Silicates.
5. Sulfides and native elements.

In addition, volatiles are present in this association.

The first three groups of minerals are the major species. They form paragenetic associations among themselves (Fig. 1). The compositions of individual minerals are variable.

Carbonates (Fig. 1A) are represented by two subgroups: (a) calcite – dolomite – magnesite (including ferromagnesite, breunnerite and sideroplesite), and (b) sodium carbonates: eitelite, nyerereite and nahcolite. The minerals form paragenetic assemblages between similar cation species: calcite with nyerereite (sample #8/103) and nahcolite (sample #8/103-2), and dolomite and magnesite with eitelite (sample #8/102).

Magnesite and dolomite, in some samples (e.g., #8/108), comprise the major part of the inclusions. The inclusion has an euhedral shape, which implies that it has the ‘negative’ morphology that is common to syngenetic mineral inclusions in diamond (Fig. 2). Some grains of dolomite are enriched with Fe and Mn, and are, possibly, ankerite (based on the observed EDX spectra).

Calcite occurs both as single, isolated crystals, 10-30 µm in size (sample #8-103; Kaminsky et al. 2009) and in association with other minerals (Table 1; Fig. 3), including CaSi-perovskite (Brenker et al. 2007) that supports its formation within the lower mantle. Some analyzed grains contain admixtures of Sr (0.74 at. %) and Ba (trace contents), confirming their primary origin. The structure of calcite in grains associated with CaSi-perovskite was proven by both Raman spectra and X-ray diffraction (Brenker et al. 2007).

Calcite, dolomite and magnesite are primary mineral phases in the lower mantle association within Juina diamond (Figs. 2 and 3). This contrasts to occurrence of these carbonate minerals in the well-studied natrocarbonatitic lavas from the Oldoinyo Lengai, Tanzania, where calcite and dolomite (magnesite is not recorded in those lavas) are considered to have formed through transformation of the primary natrocarbonatitic association minerals (e.g., Zaitsev and Keller 2006). Na-carbonates, - eitelite, nyerereite and nahcolite in both associations (from the Deep Earth and from Tanzanian volcanoes) are similar to each other, however.

The second group of minerals is represented by a variety of compounds of volatile elements (Cl, F, P and S), – **halides, fluorides, phosphates and sulfates** (Fig. 1B). Despite differences in their compositions, these minerals closely associate with one another and with the aforementioned carbonates. Among halides, in addition to most common halite, sylvite and hydrophillite, there occurs a rare mineral, cotunnite (PbCl₂). In contrast to halite and sylvite, which are typical for the Oldoinyo Lengai rocks (e.g., Zaitsev and Keller 2006), cotunnite was identified before only as a post-volcanic (fumarole) product in active volcanoes in association with other halides, or as a

secondary mineral in some lead mines. In Juina carbonatite cotunnite associates, in addition to other halides, with another rare lead mineral, plattnerite (PbO_2).

Fluorides are another major class within this mineral group. Among them, oskarssonite (AlF_3), parascandolaite (KMgF_3) and a new mineral $[(\text{Ba},\text{Sr})\text{AlF}_5]$ were identified (Kaminsky et al. 2013, 2015a). They form assemblages in close association with phosphates.

Among phosphates, in addition to common apatite, two new mineral species were identified: mixed-anion $[\text{Na}_4\text{Mg}_3(\text{PO}_4)_2(\text{P}_2\text{O}_7)]$ and Fe-diphosphate $[\text{Fe}_2\text{Fe}_5(\text{P}_2\text{O}_7)_4]$ (Kaminsky et al. 2013). The phosphates form irregular grains in dolomite and fluoride.

Anhydrite, the only sulfate in the carbonatitic association, forms assemblages with all halides, but not with the carbonates; whereas phosphates associate with both carbonates and halides, as well as with fluorides.

Oxides (Fig. 1C), which comprise the major part of the lower mantle, are subordinate in the carbonatitic association. Two major subgroups are distinguished within this class, in the studied carbonatitic associations: (a) perovskites and (b) Mg- and Fe-oxides. Perovskites are represented by CaSi-perovskite and CaTi-perovskite; to date no bridgmanite has been observed. Within the Juina inclusions, the Mg- and Fe-oxides do not form a compositional spectrum, unlike in the ultramafic association, where Mg-index of ferropericlase varies from 36 to 90. In contrast, Mg- and Fe-oxides from the carbonatitic association have only 7-15 at. % Fe and 2-15 at. % Mg, respectively (samples ##8/103 and 8/108), and may be considered as almost pure periclase and wüstite.

In addition to minerals of these two subgroups, single grains of accessory oxides, such as ilmenite, rutile with an $\alpha\text{-PbO}_2$ structure, ilmenorutile, hematite and plattnerite (PbO_2) occur. The presence of plattnerite in association with cotunnite (PbCl_2) may not be accidental. Various spinel accessory varieties (hercynite, magnesioferrite and magnetite) also occur; their role is unclear.

Of particular interest is the presence of silica. In studied specimens, SiO_2 occurs in form of coesite. However, initially it may have been stishovite or post-stishovite phases, which are unstable under decreasing pressure conditions.

Silicates also occur as single grains in the carbonatitic association. Some of them (wollastonite, monticellite and cuspidine) are secondary phases, formed as a result of the decomposition of CaSi-perovskite during the ascent of the host diamond to the surface (Kaminsky et al. 2014). A mineral phase with a composition that of olivine has been identified not only in the carbonatitic association, but within the ultramafic one as well (Kaminsky 2012). It has yet to be studied structurally, and its nature remains unclear.

The other minerals, occurring in the carbonatitic association, - **sulfides** (pentlandite, violarite and others) **and native iron** belong to the matrix of the Deep Earth. Native iron contains 3.5 at. % Ni (sample #8/102; Kaminsky et al. 2013) and, by that, is similar to meteoritic kamacite. These minerals should be considered as accessory phases.

In general, the Juina carbonatite association is predominantly carbonate (natrocarbonate) – halide – fluoride in composition with a minor (accessory) admixture of oxides, silicates and other minerals. It is close, by mineral composition, to the natrocarbonatitic association.

Natrocarbonatitic associations

Four major types of carbonatite are known: calcite carbonatite, dolomite carbonatite, ferrocarbonatite, and natrocarbonatite (International ... 1989). While the first three types are distributed widely, natrocarbonatites were known, until recently, only in one location: - as recent lavas in the Oldoinyo Lengai volcano in Tanzania ('lengaites'; Kresten 1983), and within the Kerimasi and Tinderet carbonatites (Mitchell 2005; Zaitsev et al. 2014). Natrocarbonatite rocks are composed mainly of nyerereite $\text{Na}_2\text{Ca}(\text{CO}_3)_2$ and gregoryite $(\text{Na},\text{K},\text{Ca})_2\text{CO}_3$, with minor sylvite and fluorite in the groundmass. As such, these demonstrate comparable mineralogical and bulk chemical (Na-Cl-F) features to the carbonatitic association identified in lower-mantle Juina diamonds. In addition, the Juina carbonatitic association is enriched in REE, Sr and Nb, which is common for primary carbonatites. This demonstrates that the two associations, lengaite and Juina carbonatite may be considered as representatives of the same, natrocarbonatitic type.

There are, however, some differences in the compositions of the two associations. For example, in addition to Na-carbonates, calcite, dolomite and magnesite also occur in Juina carbonatite entrapped in diamond. Further, calcite and dolomite in the Juina carbonatitic association represent primary crystallizing phases, while in lengaites this mineral association is the result of transformation of a 'primary' natrocarbonatitic association (Zaitsev and Keller 2006). Fluorite was not identified in Juina samples, while other fluorides, observed in Juina, are not characteristic for lengaites. Such differences may be the result of different source media compositions, their significantly differing depths of origin, and paragenesis (see below).

Depth of origin of the Juina diamond hosted carbonatites, and the stability of carbonates under lower-mantle conditions

Minerals comprising the deep mantle natrocarbonatitic association are stable within a wide range of P-T conditions. With the exception of secondary minerals, such as monticellite, wollastonite, etc., they (or their polymorphs) are stable under conditions found within the lower-mantle.

Carbonates are major rock-forming minerals in the association. Close assemblages of carbonates with lower-mantle oxides as inclusions in diamonds from the Juina area have been identified, and include: calcite with CaSi-perovskite and CaTiO₃ (Brenker et al. 2007), and calcite, dolomite and nyerereite with periclase and wüstite (Kaminsky et al. 2009). The relationships between these phases and associations support an origin for the Juina natrocarbonatitic association within the lower mantle, i.e., below the 660-km discontinuity. These data agree with both experimental and theoretical data on the stability of carbonates in the Deep Earth.

Magnesite is considered as the primary carbonate phase within the Deep Earth (Brenker et al. 2007; Lin et al. 2012; Scott et al. 2013); it is the most stable carbonate under conditions prevalent in the lower-mantle. While it has been suggested that magnesite will decompose into MgO and CO₂ at extreme pressures (Irving and Wyllie 1973) or, in association with SiO₂, into ferropericlase, with the carbon forming diamond (e.g., Seto et al. 2008), both experimental and theoretical studies have demonstrated that it is stable under pressure conditions of up to 113-115 GPa (~2,500 km depth) and $T = 2,100-2,200$ K, when magnesite *R-3c* structure transforms into a pyroxene-type orthorhombic MgCO₃ ('magnesite II', Isshiki et al. 2004; Skorodumova et al. 2005; Oganov et al. 2006, 2008; Panero and Kabbes 2008), and then, at ~200 GPa from the pyroxene-structure into the CaTiO₃-structure (Skorodumova et al. 2005). Under such extreme pressure conditions only at very high temperatures (> 2,700 K) would magnesite decompose, with the formation of MgO and diamond (Solopova et al. 2014).

The calculated stability of CaCO₃ reaches ~80 GPa pressure and $T = 3,300$ K (Ivanov and Deutsch 2002). Within a pressure range of 1-5 GPa and $T = 800-2,000$ K calcite changes its rhombohedral *R-3c* structure to that of rhombohedral aragonite (Irving and Wyllie 1973; Ivanov and Deutsch 2002), although some triclinic (*P-1*) modifications are stable at pressures of up to 30 GPa (Merlini et al. 2014). At pressures near 50 GPa the aragonite orthorhombic structure undergoes a transition to a trigonal, post-aragonite structure (Santillán and Williams 2004a), and after reaching 137 GPa (i.e., at the core/mantle boundary) the pyroxene-type *C222₁* structure of CaCO₃ may be expected (Oganov et al. 2006; Ono et al. 2007).

However, the grain from our sample #RS-35, where calcite associates with CaSi-perovskite, was identified not only by Raman spectrum, but by X-ray diffraction as well and is demonstrated to have the calcite structure (Brenker et al. 2007). This potentially indicates, that the calcite rhombohedral structure can be stable, in the natural environment, under pressure conditions of up to at least 14-24 GPa (Shim et al. 2000). In our other studied samples, calcite was identified by analogy on the basis of its composition only; aragonite orthorhombic or trigonal structure of some of those grains may not be excluded.

Dolomite, in our sample #8/102, forms a syngenetic assemblage with magnesite (Fig. 2; Kaminsky et al. 2013). In another sample (#8/108), a spherical inclusion of wüstite + periclase in dolomite was identified, pointing to its formation under pressure conditions exceeding 86 GPa (Kaminsky et al. 2015a). This agrees with the experimental data of Mao et al. (2011), who demonstrated, that CaMg(CO₃)₂ is stable under lower-mantle conditions at pressures of up to 83 GPa and, possibly, higher, and contrasts earlier data on the decomposing of dolomite into

magnesite + aragonite, under lower-mantle P-T conditions (Biellmann et al. 1993; Luth 2001; Shirasaka et al. 2002). Recent experiments confirm that high-pressure polymorphism in dolomite could stabilize $\text{CaMg}(\text{CO}_3)_2$; this composition transforms at ~17 GPa into ‘dolomite-II’ (with a monoclinic structure, according to Santillán et al. 2003, or an orthorhombic structure, according to Mao et al. 2011, or a triclinic structure for Fe-dolomite, according to Merlini et al. 2012) and then, at ~35-41 GPa, into ‘dolomite-III’ (with monoclinic structure, according to Mao et al. 2011 or a triclinic structure in the case of Fe-dolomite, according to Merlini et al. 2012).

Volume differences between the three major carbonates: CaCO_3 , MgCO_3 and $\text{CaMg}(\text{CO}_3)_2$ under high pressures are minimal, implying that energetic differences between these phases are small (Santillán et al. 2003) and they all may be present in the lower mantle, depending on the chemical composition of the media, or within the same association under variable conditions. Among carbonates, the species with small radii of divalent cations (less than 1 Å, i.e., Mg^{2+} with 0.89 Å and Fe^{2+} with 0.78 Å radii) are more stable, while carbonates with large radii (Ca^{2+} with 1.12 Å radius) are less stable (Santillán and Williams 2004b). However, real mantle composition depends, first of all, on the chemical composition of the media; and CaCO_3 and $\text{CaMg}(\text{CO}_3)_2$ may both be predominant mineral phases if the media is enriched in Ca. Rhombohedral *R-3c*-structured carbonates are the most stable under high pressure conditions (Santillán et al. 2005); and the boundary of the calcite-I to aragonite phase transition may be of up to 44 GPa, as determined from CaCO_3 and CdCO_3 (Liu and Lin 1997; Santillán and Williams 2004b, Fig. 10).

Other minerals from the studied natrocarbonatitic association are experimentally proven to be stable at ultrahigh pressure conditions as well: eitelite ≥ 21 GPa (Kiseeva et al. 2013), parascandolaite ≥ 50 GPa (Aguado et al. 2008).

Of particular interest are finds, in the Juina natrocarbonatitic association, of almost pure periclase and wüstite. In sample #8/103, two mineral species occur as coexisting, euhedral nanoinclusions in a porous carbonate matrix: periclase with $mg = 0.85-0.92$ and wüstite with $mg = 0.02-0.15$; no ferropericlase or magnesiowüstite of intermediate composition (which is characteristic for the oxide lower mantle) were observed (Kaminsky et al. 2009). Recently, in another sample #8/108, almost pure periclase and wüstite (with admixture of 5 wt. % Fe and Mg, respectively) were observed, forming a spherical-shaped inclusion in a porous dolomite-calcite matrix: the core is composed of wüstite, and the rim is composed of periclase, both with spinel-type structures (Fig. 4) (Kaminsky et al. 2015a).

It is proven experimentally, that ferropericlase dissociates into almost pure wüstite and an Mg-rich phase at 86 GPa and 1000 K (Dubrovinsky et al. 2000). As such, the natrocarbonatitic association in the Deep Earth formed at depth conditions corresponding or exceeding 86 GPa, corresponding to a depth in excess of 2,000 km. In other samples from Juina, iron-rich (Mg,Fe)O inclusions in diamond have been attributed, apparently, to the lowermost mantle (Wirth et al. 2014).

A recent detailed study of magnesiowüstite of composition $\sim(\text{Mg}_{0.35}\text{Fe}_{0.65})\text{O}$, included in another diamond from the Juina area, led to the conclusion over its origin in one of the ultra-low velocity zones at the base of the mantle, and with this reaching the surface through diapiric upwelling culminating in kimberlite eruption (Wirth et al. 2014).

A major factor of the stability of carbonate minerals and, hence, the formation of the carbonatite association in the Deep Earth is oxygen fugacity. Recent thermodynamic calculations have shown that typical values of oxygen fugacity in zones of diamond formation in the lower mantle lie between the iron-wüstite buffer and six logarithmic units above this level (Ryabchikov and Kaminsky 2013). These calculations are proven by the formation of ferropericlase in the lower-mantle within a wide range of $\Delta\log f_{\text{O}_2}$ (IW), from -1.76 to 7.86 (Kaminsky et al. 2015b). This observation makes formation of the carbonatitic association within the lower mantle realistic.

Genesis of the natrocarbonatitic association in the Deep Earth

Carbon, diamond and hotspots

In numerous works (e.g., Mao et al. 2011 and references therein) the presence of carbonate species in the lower mantle has been considered a result of the subduction of lithospheric plates and entrained sedimentary carbonates into the lower mantle. An alternative explanation, however, suggests the origin of carbonate minerals in the lower mantle may instead be juvenile carbon. Deep Earth's interior is a primary reservoir for this element; it has been calculated that more than 90 % of Earth's carbon is stored deep within this reservoir (e.g., Javoy 1997; McDonough 2003). The Earth's core, where the carbon content is suggested at ~ 5 wt. %, supplies carbon to the mantle over geological time via the mechanism of grain boundary diffusion that results in the formation of C-H-O volatiles (e.g., Hayden and Watson 2008). As a result, the Earth's mantle is the largest Earth's reservoir of carbon (Dasgupta and Hirschmann 2010) with an average C concentration of 100-120 ppm (Zhang and Zindler 1993; McDonough 2003).

It has been supposed, on the basis of a series of petrologic and isotopic evidence, that carbonatite magmas may originate in the deepest parts of the mantle (Dalou et al. 2009; Bell and Simonetti 2010; Collerson et al. 2010) and be linked to the production of large igneous provinces (Ernst and Bell 2010). Our data on the existence of carbonatitic inclusions in lower-mantle diamond support these ideas. We suggest that the formation of carbonatite and superdeep diamond initiated within the lower-mantle, in regions where hotspot/ plumes occur. The observed presence of diamond in modern, Hawaiian plume products, where these occur in a low-degree melt glass, rich in C, Cl, and comprising an exsolved, predominant CO₂, gas-fluid phase (Wirth and Rocholl 2003) gives a basis for such a proposition. Moreover, recent find of carbonatitic and iron carbide-containing inclusion within the same diamond grain (Kaminsky et al., 2015) confirms the suggestion about the formation of carbonatitic liquid in the deepest parts of the mantle.

'Primary', deep hotspot plumes (Courtillot et al. 2003) are thought to originate at the thermal boundary layer at the base of the lower mantle (e.g., Zhong 2006; Boschi et al., 2007). In some areas of the core-mantle boundary (CMB), anomalous seismic properties within thin zones (~ 10 km thick and 50-100 km wide) were identified as Ultra-Low Velocity Zones (ULVZ) due to their drop in seismic velocities greater than 10 % relative to the background mantle. They are considered as zones of partial melting, which may result from vigorous, small-scale convection or instabilities in the thermal boundary layer at the base of the mantle (e.g., Wen and Helmberger 1998; McNamara et al. 2010). ULVZ material can become entrained in mantle plumes (McNamara et al. 2010).

Formation of carbonatitic partial melts in the lowermost mantle

The adiabatic *temperature profile in the mantle*, based on experiments and the use of the equation of state of MgO from various sources, was calculated by Katsura et al. (2010). The adiabatic temperature coefficient in the lower mantle was found to be 0.3 K/km (lower than in the upper mantle), and the adiabatic temperature at a depth of 2,700 km was estimated at $2,730 \pm 50$ K, if convective heat still dominates in this region. However, the lower mantle is not homogeneous seismically (Bunge et al. 2001), and 3D spherical convection is not applicable to the entire Deep Earth. There are data, based on the elasticity of silicate perovskite, that the real temperature profile in the lower mantle is much steeper than the average mantle adiabat (AMA), and below 1,500 km the actual geotherm is super-adiabatic (da Silva et al. 2000). The computer simulations predict non-adiabatic excess of 100-300 degrees for the lower mantle because a significant proportion of mantle heat source is internal (Bunge et al. 2001). Calculations based on measurement of perovskite shear modulus also demonstrate that lower mantle seismic properties may not match an adiabatic geotherm; geotherms may have larger temperature gradients (from 0.5 to 0.9 K/km between 800 and 2,700 km), and indeed the temperature may reach 3,400 K at a depth of 2,700 km (Matas et al. 2007).

The pyrolite *solidus temperature* at the core-mantle boundary (CMB) was advocated to be near that of the core temperature (~ 3,800-4,000 K), suggesting partial melting of the lower mantle possible (Zerr et al. 1998). More recent experimental data on the solidus temperature at the core-mantle boundary (135 GPa pressure) are even higher: $4,180 \pm 150$ K for fertile peridotite (Fiquet et al. 2010) and $4,150 \pm 150$ K for the chondritic composition (Andrault et al. 2011). These data are based on a study of both pyrolitic and chondritic solidi as 'dry' systems composed of only major elements (Si, Al, Fe, Ca and Mg), while the presence of alkalis and particularly C, O, H and other volatiles depress the solidus position drastically. The experimental data performed at 10-15 GPa pressure conditions demonstrated that the presence of C-O-H fluid and/or even the minor admixture of alkalis in peridotite

produce carbonatitic melt at temperatures lower than adiabatic ones by as much as 400-500 °C (Litasov et al. 2013a, 2013b). It was demonstrated, in experiments at up to 80 GPa, that the melting behavior of simple carbonate systems is a suitable proxy for many carbonate-bearing lithologies (Thomson et al. 2014).

In this case even a slight increase of temperature initiates the process of *partial melting*, which is observed in ULVZs at the CMB by P- and S-wave velocity reductions (Lay et al. 2004 and references therein). Such areas of partial melting are likely the roots to mantle hotspots. These hotspots are believed to be caused by the presence of chemical heterogeneities with high concentrations of fusible elements (Andrault et al. 2011). It has been calculated theoretically (Wyllie and Ryabchikov 2000), and proven experimentally (Dalton and Presnall 1998), that low-fraction melts generated near the solidus of carbon-containing mantle lherzolite at high enough oxygen fugacity values should have a carbonatite composition, with CO₂ contents of ~ 45 wt.%.

Given that estimates of the average concentrations of carbon and phosphorus in mantle peridotite are closely similar (Palme and O'Neill, 2003), the *near-solidus melts should have a carbonate-phosphate composition* (Ryabchikov and Hamilton, 1993a, 1993b) - exactly as we have observed in the diamond-hosted, Juina carbonatite association. In experiments, low-fraction volatile-rich carbonatitic melts extract from parental peridotite many incompatible elements (Ryabchikov et al. 1991). The natural carbonatite association is rich in chlorine, fluorine and other elements, which form, in addition to carbonates and phosphates, fluorides and chlorides (Kaminsky et al. 2009, 2013).

As a result, carbonatitic melts can be produced from volatile- and/or alkali-rich peridotite (chondrite) at the base of the lower mantle, in the roots to mantle hotspots at the CMB.

Evolution of carbonatitic melts in the lower mantle

Our suggested model for the formation and evolution of carbonatite melts in the Deep Earth is illustrated in Fig. 5.

The *first, super-adiabatic, stage A* occurs within the lowermost mantle, starting from the CMB (D'' layer). A local increase in the thermal condition, caused most likely by processes in the upper core, triggers low-degree partial melting of lower-mantle material, enriched in carbon and other volatiles. Within chambers, approximately 10-20 km thick and 50-100 km wide, carbonatitic melts containing ≥ 45 wt. % CO₂ are formed. Such chambers, which are roots to future hotspots, begin to migrate upwards creating local 3D convection. Partial carbonatitic melts comprise 1-2 vol. % of the mantle chamber material, and are strongly enriched in volatiles. These partial melts infiltrate mantle lithologies and percolate into regions of the mantle. Carbonate melts have been shown to be highly mobile; they can travel upward into and through the upper mantle at velocities exceeding hundreds to thousands of meters on time scales of 0.1–1 m.y.; i.e., several orders of magnitude higher than those previously found for basaltic liquids (Hammouda and Laporte 2000; Kono et al. 2014).

During the *second stage B* partial melts develop at near-adiabate conditions. During this stage oxide phases can be crystallized; some of them may form immiscible wüstite-periclase associations. The temperature conditions within this stage correspond to multicomponent, completely miscible carbonate liquid system (Litvin et al. 2014). According to experimental data, as temperature falls, the invariant eutectic assemblage Mg-Fe-magnesite (Mg,Fe)CO₃ + Na-carbonate Na₂(Mg,Fe)(CO₃)₂ + liquid is formed (Litvin et al. 2014).

At lower P-T parameters, the initial carbonatitic system dissolves into two parts. Experiments at 5-16.5 GPa and high temperature demonstrated the presence of two immiscible carbonatitic systems: carbonate-silicate and chloride-carbonate (Litasov and Ohtani 2009; Safonov et al. 2011). The latter one may be considered as an analogue to the natrocarbonatitic Juina lower-mantle association, while carbonate-silicate association has yet to be observed in mantle conditions. At high temperatures the two immiscible liquids tend to coexist (Litasov and Ohtani 2009); however, the temperature for this immiscibility is not yet known. One may suggest that it occurs within the P-T conditions of the near-adiabatic stage B, where two immiscible carbonatitic melts separate. To date, only the chloride-carbonate (natrocarbonatitic) association is known, however, there is no reason to exclude the existence of the second one in the Deep Earth.

Most likely, during the near-adiabatic stage B, formation of the 'ultra-deep' diamond occurs. Carbonatite media is favorable to diamond formation. It has been established, experimentally that a concentration barrier of

diamond nucleation (CBDN) arises at a particular ratio of carbonate to silicate components (Litvin 2009). Studies of different systems have revealed a dependence of barrier position on the chemical composition of the system and the inhibiting effect of silicate components on the nucleation density and rate of diamond crystal growth. In multicomponent silicate-carbonatite solvent, the CBDN is within the range of carbonatite compositions (< 50 wt. % silicates). Diamonds have been experimentally crystallized in melts of the lower mantle diamond parental carbonate - magnesiowüstite – bridgmanite – carbon system, at conditions found within the upper part of the lower-mantle (20-30 GPa) (Litvin et al. 2014). It is apparent that admixtures of P, Cl and F within carbonatitic melt are of great importance in the formation of diamond (Palyanov et al. 2007; Sonin et al. 2008). Within both fibrous and monocrystalline upper-mantle diamonds, carbonatitic and saline high-density fluids (HDF) were identified, some of them are purely carbonatitic in composition (e.g., Schrauder and Navon 1994; Izraeli et al. 2001; Klein-BenDavid et al. 2006, 2009; Weiss et al. 2014). They are considered as relics of diamond-forming media. The formation of lower-mantle diamonds may have happened either from carbonatitic melt or from high-density fluids.

The *third stage C* occurs at sub-adiabatic conditions. Various carbonate phases (CaCO_3 , MgCO_3 , $\text{CaMg}(\text{CO}_3)_2$ and others) form within this stage, and major phase transformations occur in subsolidus conditions, with the participation of remnant fluid phases. In experiments, the assemblage Mg-Fe-magnesite $(\text{Mg,Fe})\text{CO}_3$ + Na-carbonate $\text{Na}_2(\text{Mg,Fe})(\text{CO}_3)_2$, with Na_2CO_3 solidification at the lowermost isobaric temperature 1300 K (Litvin et al. 2014) was observed.

At or near to the Earth's surface diamonds with carbonatitic inclusions are delivered, as xenocrysts, by kimberlitic volcanism. The kimberlite magmas form at shallower depth, as a result of partial melting of peridotite within the upper mantle and/or transition zone. The diamonds entrained by kimberlitic magma en route to the surface are formed both in the lower-mantle and upper-mantle/transition zone (dominating) and, as such, contain inclusion suites of both carbonatitic and ultramafic mineral associations.

Conclusions

1. In addition to ultramafic and mafic (an analog to eclogite) associations, a primary natrocarbonatitic association occurs in the lower mantle. This latter association is hosted within inclusions in kimberlitic diamond from the Juina area, Mato Grosso State, Brazil.
2. The initial lower-mantle carbonatitic melt formed as a result of low-fraction partial melting of lower-mantle material at the core-mantle boundary. It was rich in P, F, Cl and other volatile elements.
3. During ascent to the surface the initial carbonatitic melt dissociated into two immiscible parts, a carbonate-silicate and a chloride-carbonate melt. The latter melt is parental to the observed natrocarbonatitic lower-mantle association.
4. Diamonds with carbonatitic inclusions were formed in carbonatitic melts or high-density natrocarbonatitic fluids.

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Tables

Table 1. List of minerals of the primary carbonatitic association in the Deep Earth.

Mineral	Index	Formula	Association	Sample # (Reference)
Carbonates				
Calcite	Cc	CaCO ₃	Phl	(1)
			CaSiPvk+CaTiPvk+Ol	(2)
			Mont+Wo Il+Cusp	8/104-1 (3)
			Per+Wu	8/103 (4)
			Sp+Ap	
			Sp+Phl(?)	
			Nyer+Wo Il+CaGa	
			Nah +Phl	8/103-2 (4)
Dolomite	Dol	CaMg(CO ₃) ₂	Dol+Ap+Sp+Sulf+Per+Wu+Phl(?)	8/108 (6)
			Ilm	8/106 (3)
			Sp+Phl	
			Cc+Ap+Sp+Sulf+Per+Wu+Phl(?)	8/108 (6)
Magnesite	Mgs	Mg(CO ₃)	Mgs+Eit+Ha+Sy+Phl+Pent+Vi+Sp+P1	8/102 (5)
			FeMgs+ Fe ⁰	
Fe-magnesite	FeMgs	(Mg,Fe)(CO ₃)	Mgs+Fe ⁰	8/102 (5)
Eitelite	Eit	Na ₂ Mg(CO ₃) ₂	Dol+Mgs+Ha+Sy+Phl+Pent+Vi+Sp+P1	8/102 (5)
Nyerereite	Nyer	(Na,K) ₂ Ca(CO ₃) ₂	Cc+Wo Il+CaGa+Ap+Wu(?)	8/103 (4)
			Ol+Sp+Ap+Phl(?)	
Nahcolite	Nah	NaHCO ₃	Cc+ Phl	8/103-2 (4)
Halides				
Halite	Ha	NaCl	Coes+Sy+Hy+Cot+Plat+TiO ₂ +Anh	8/105 (3)
			Dol+Mgs+Eit+Sy+Phl+Pent+Sp+P1	8/102 (6)
			No	8/102 (6)
Sylvite	Sy	KCl	Coes+Ha+Hy+Cot+Plat+TiO ₂ +Anh	8/105 (3)
			Dol+Mgs+Eit+Ha+Phl+Pent+Vi+Sp+P1	8/102 (5)
Hydrophilite	Hy	CaCl ₂	Coes+Ha +Sy +Cot+Plat+TiO ₂ +Anh	8/105 (3)
Cotunnite	Cot	PbCl ₂	Coes+Ha +Sy +Hy+Plat+TiO ₂ +Anh	8/105 (3)
Fluorides				
Oskarssonite	Osk	AlF ₃	P2+BaFl	8/102 (5)
			SiO ₂ +Fe-O+Mil Hem+ SiO ₂	
Parascandolaite	Prs	KMgF ₃	Hc+OrthMgO	8/108 (6)
New (no name)	BaFl	(Ba,Sr)AlF ₅	P2+Osk	8/102 (5)
Sulfates				
Anhydrite	Anh	CaSO ₄	Coes+Ha+Sy+Hy+Cot+Plat+TiO ₂	8/105 (3)
Phosphates				
Apatite	Ap	Ca ₅ [PO ₄] ₃ (F,Cl)	Nyer+Ol+Sp+Phl(?)	8/103 (4)
			Nyer+Cc+Wo Il+Wu(?)	
			Cc+Sp+Wu	
New (no name)	P1	Na ₄ Mg ₃ (PO ₄) ₂ (P ₂ O ₇)	Cc+Dol+Sp+Sulf+ Per+Wu+Phl(?)	8/108 (6)
New (no name)	P2	Fe ₂ Fe ₅ (P ₂ O ₇) ₄	Dol+Mgs+Eit+Ha+Sy+Phl+Pent+Vi+Sp	8/102 (5)
Osk+BaFl				8/102 (5)
Oxides				
CaSi-perovskite	CaSiPvk	CaSiO ₃	Cc+Ol+CaTiPvk	(2)
CaTi-perovskite	CaTiPvk	CaTiO ₃	Cc+Ol+CaSiPvk	(2)
Periclase	Per	MgO	Cc+Wu	8/103 (4)
			Dol+Ca+Ap+Sp+Sulf+ Wu+Phl(?)	8/108 (6)
Orrthorhombic MgO	OrthMgO	MgO	Parasc+Hc	8/108 (6)
Wüstite	Wu	FeO	Cc+Per	8/103 (4)
			Cc+Sp+Ap	
			Nyer+Cc+Ap+Wo Il	
Dol+Ca+Ap+Sp+Sulf+Per+ Phl(?)	8/108 (6)			
Coesite	Coes	SiO ₂	Ha+Sy+Hy+Cot+Plat+TiO ₂ +Anh	8/105 (3)

Non-crystalline		SiO ₂	Osk+Fe-O+Mil	8/102 (5)
Ilmenite	Ilm	FeTiO ₃	Dol Sp+Phl	8/106 (3)
Rutile(?) (α-PbO ₂ structure)	Ru(?)	TiO ₂	Coes+Ha+Sy+Hy+Cot+Plat+ Anh	8/105 (3)
Ilmenorutile(?)	IlmRu(?)	(Ti,Nb,Fe)O ₂	Osk+Hem+SiO ₂	8/102 (5)
Hematite	Hem	Fe ₂ O ₃	Osk+ SiO ₂	8/102 (5)
Plattnerite	Plat	PbO ₂	Coes+Ha+Sy+Hy+Cot+Ru+Anh	8/105 (3)
Spinel-hercynite	Hc	FeAl ₂ O ₄	Parasc+OrthMgO	8/108 (6)
Spinel-magnesioferrite	Sp	(Mg,Fe)(Al,Fe) ₂ O ₄	Dol+Phl+Ilm Nyer+Ol+Ap+Phl(?) Cc+Phl(?)	8/106 (3) 8/103 (4)
Spinel-magnetite	Sp	Fe ₃ O ₄	Cc+Ap	8/103 (4)
Spinel non-specified	Sp	Mg-Fe-Al-O	Dol+Ca+Ap+ Sulf+ Per+Wu+Phl(?)	8/108 (6)
Silicates				
"Olivine"	Ol	(Mg,Fe) ₂ SiO ₄	Cc+CaSiO ₃ +CaTiO ₃ Nyer+Sp+Ap+Phl(?)	(2) 8/103 (4)
Wollastonite-II (high)	Wo II	CaSiO ₃	Cc+Cusp+Mont Cc+Nyer+CaGa+Ap+Wu	8/104-1 (3) 8/103 (4)
Monticellite	Mont	CaMgSiO ₄	Cc+Wo II+Cusp	8/104-1 (3)
Cuspidine	Cusp	Ca ₄ [Si ₂ O ₇](F,OH) ₂	Cc+Wo II+Mont	8/104-1 (3)
Ca-Garnet	CaGa	Ca ₃ (Fe,Zr,Ti) ₂ [(Si,Al)O ₄] ₃	Cc+Nyer+Wo II+Ap+Wu	8/103 (4)
Phlogopite	Phl	KMg ₃ [Si ₃ AlO ₁₀](F,OH) ₂	Cc Dol+Ilm+Sp Nyer+Ol+Sp+Ap Cc+Sp Nah+Cc Dol+Mgs+Eit+Ha+Sy+Pent+Vi+Sp+P1 Dol+Ca+Ap+Sp+Sulf+ Per+Wu	(1) 8/106 (3) 8/103 (4) 8/103-2 (4) 8/102 (5) 8/108 (6)
Sulphides				
Pentlandite	Pent	(Fe,Ni) ₉ S ₈	Dol+Mgs+Eit+Ha+Sy+Phl+Vi+Sp+P1	8/102 (5)
Violarite	Vi	FeNi ₂ S ₄	Dol+Mgs+Eit+Ha+Sy+Phl+Pent+Sp+P1	
Millerite	Mil	NiS	Osk+SiO ₂ +Fe-O	8/102 (5)
Fe(-Ni) Sulfide	Sulf	Fe-Ni-S	Dol+Ca+Ap+Sp+Per+ Wu+Phl(?)	8/108 (6)
Fe-Ni-Cu Sulfide	Sulf	Fe-Ni-Cu-S	(Cc+Wo II+Cusp+Mont)	8/104-2 (3)
Native elements				
Ni-iron	Fe ⁰	Fe-Ni	Mag+FeMgs	8/102 (5)

Fig. 2. Syngenetic inclusion of dolomite and magnesite in diamond. Sample #8/102, foil #3327. TEM image. Scale bar 1 μm .

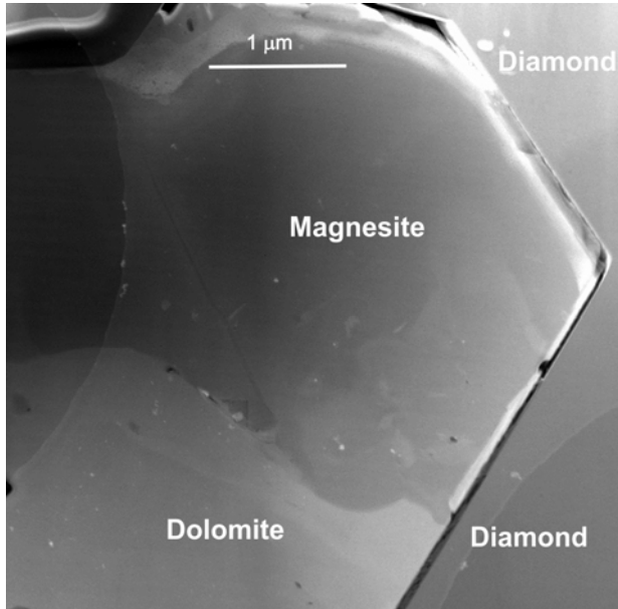


Fig. 3. Calcite and nyerereite in carbonatitic inclusion in diamond. Sample #8/103, foil #1734. TEM image. Scale bar 100 nm.

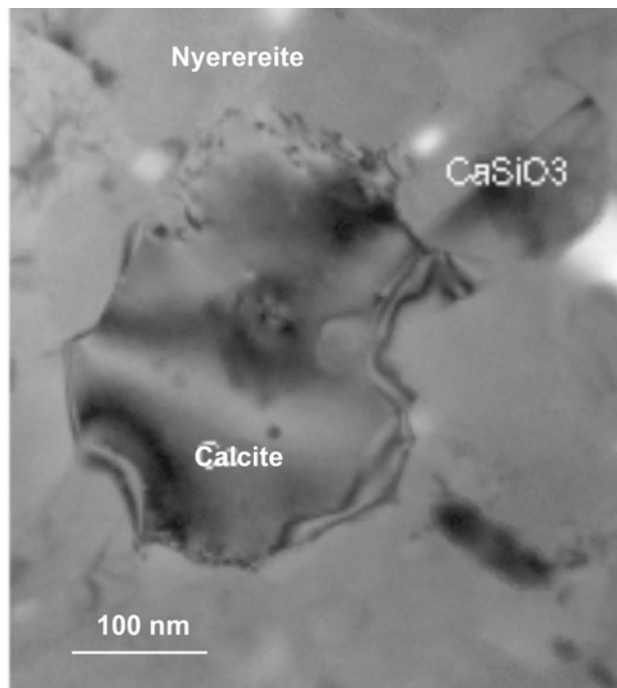


Fig. 4. Spherical inclusion of wüstite (core) + periclase (rim) in dolomite matrix. Irregular dark areas are pores. Elemental map. Scale bar is 500 nm. Sample #8/108; foil #3709.

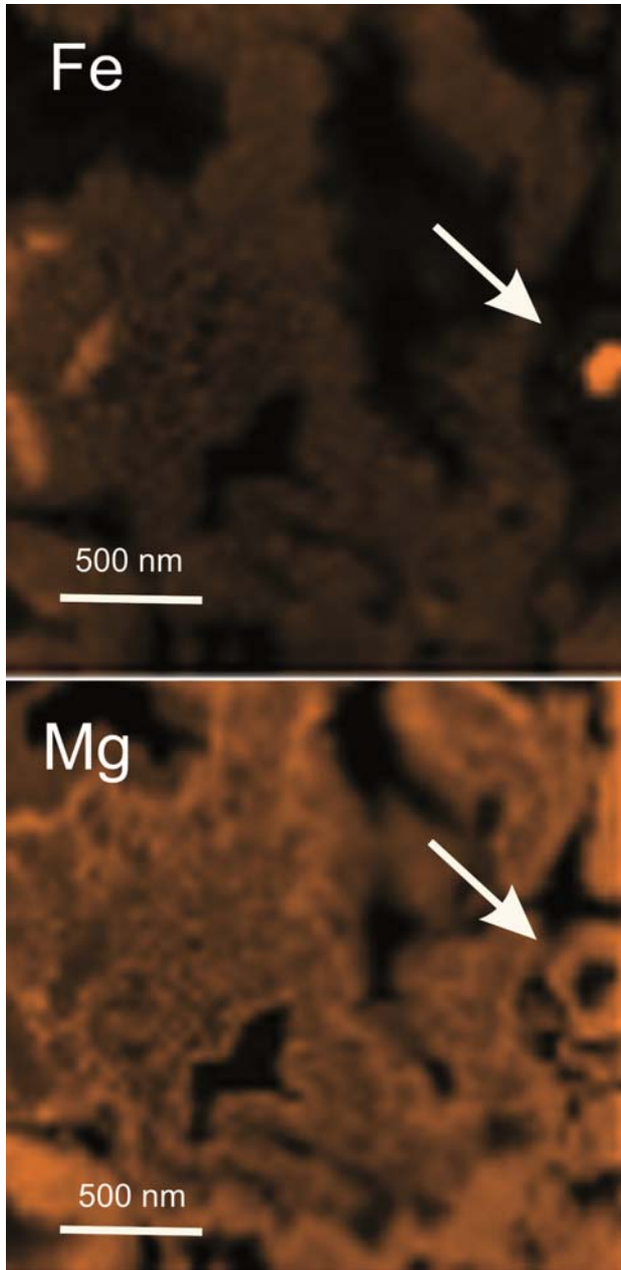


Fig. 5. A proposed model for the formation and development of the carbonatitic association in the Deep Earth.

