Modern Concepts on Diamond Genesis

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Abstract—The best-known, most well-studied diamondiferous rocks are kimberlites and lamproites. Diamonds are also found in impactites, metamorphic rocks, ophiolites, and modern volcanic rocks. Diamonds from these rocks differ from kimberlitic diamonds in size, morphology, trace-element and isotope composition, and physical properties. Differences in these characteristics are related to their different mechanisms of origin. In some cases, diamonds can be formed in “metastable” conditions under disequilibrium thermodynamic parameters, supporting the conclusion that diamond is a polygenetic mineral, formed in nature under different physicochemical and geodynamic conditions. According to thermodynamic considerations and calculations, “metastable” crystallization of diamond is mainly controlled by the size of the forming crystallites. The main effectors in decreasing the energetic barrier for nanosized diamonds are surface tension and related surface energy.

Keywords: diamond, genesis, kimberlites, lamproites, ophiolites, chromitites, impactites, phase transitions

INTRODUCTION

For several decades, monographs “Diamond Genesis” (Williams, 1932) and “Diamond Fields of South Africa (Wagner, 1914) have been the main sources of data on diamond geology and the genesis of diamonds. Later publications were dedicated to diamonds from placer deposits in India, Indonesia, Southeast Australia, Brazil, and Urals. Diamonds were well known to be found in these deposits before the discovery of South African diamonds, but were thought to be unrelated to kimberlites. Several reviews that have considered these data have been published in the USSR (Sobolev, 1951; Kukharenko, 1955; Trofimov, 1980) and France (Bardet, 1973, 1974, 1977). Throughout 1950 s to the 1990s, diamondiferous kimberlites were found on all continents, including practically all ancient cratons in Siberia, North America, Australia, China, and Brazil (Fig. 1). Studies have shown that diamonds in kimberlites are formed at large mantle depths under high temperatures and pressures. Concepts regarding diamond genesis based on existing research have been summarized in a publication by V.S. Sobolev (1960), which has served as a basis for further scientific developments for many years.

Of special significance were studies of deep-seated mineral inclusions entrapped during diamond formation, which represent means of probing parental mantle rocks and provide insights into conditions during diamond formation (Sobolev, 1974; Meyer, 1987). It has been established that diamonds can be formed in the upper mantle at depths greater than 150 km. Together with data on xenoliths in alkaline basalts, this knowledge has made it possible for the first time to study upper mantle materials, leading to the conclusion that primary diamond sources are not kimberlites but diamondiferous peridotites and eclogites, which comprise deep mantle horizons, and are destructed to supply diamonds to kimberlites originating in these rocks.

Geochemical studies, first of all, determination of carbon isotope composition in diamonds, are also of great significance. Single measurements conducted by Craig, Wickman, and Vinogradov have been continued and developed in a detailed study by Galimov et al. (1978). Galimov et al. revealed, for the first time, carbon isotope zoning and wide variations in carbon isotope composition in diamond crystals (Galimov, 1984). Studies of carbon isotope composition in diamond is a powerful tool for studies of diamond genesis (Sobolev et al., 1979; Galimov, 1984).

In the next stage of research, diamonds were discovered in a wide variety of non-kimberlitic igneous rocks, including alkaline basalts, picrites, and ultrabasic lamprophyres. The large number of these finds, including in situ in different rocks, indicates that this is a natural phenomenon (Kaminsky, 1984). High economic diamond deposits have been discovered in lamproites of the Argyle and Ellendale mines in Western Australia (Jaques et al., 1986). Some diamond deposits in North America and India have also been
found in lamproites (Scott-Smith, 1989). These discoveries significantly expanded promising territories for diamond prospecting, since diamondiferous lamproites and other rocks occur not only within ancient cratons (as do kimberlites) but also in ancient fold belts. At present, dozens of diamond finds have occurred in non-kimberlitic volcanic rocks (Kaminsky, 2007). Diamonds from both non-kimberlitic rocks and mineral inclusions within them seem to be identical to those of kimberlites, and the mechanism of their formation appears to be similar: through disaggregation of mantle ultramafic rocks during formation of lamproites and other non-kimberlitic rocks. Both kimberlites and other volcanic rocks are only diamond carriers.

During this period, diamonds (sometimes in economic concentrations) have also been discovered in rocks sharply differing from kimberlites in composition and genesis, including impactites, metamorphic rocks, ophiolites, and modern volcanic eruptions. Impactites from some astroblemes (Masaitis et al., 1972; Masaitis, 1998, 2013) and metamorphic rocks (Dobrzhinetskaya, 2012) have very high diamond concentrations, an order of magnitude higher than those in kimberlites. Diamonds in chromitites were discovered for the first time in 1980s in China (Fang and Bai, 1981) and are currently found in many ophiolite belts (Yang et al., 2021). Recently, diamonds have been found in the products of modern volcanic eruptions in Kamchatka (Gordeev et al., 2014; Karpov et al., 2014; Galimov et al., 2020). Except for impact diamonds originating from meteorite impacts, the genesis of other diamond types (metamorphic, ophiolite, volcanic) remains controversial. Regardless of exact mechanism, high-pressure diamond crystallization in the mantle is not the only mechanism of diamond formation. Diamond is a polygenetic mineral, and studies of all possible mechanisms and settings underlying its formation are of great scientific and practical significance.

It is impossible to cover in detail all modern concepts on the diamond genesis in the framework of a single journal paper. Our work thus focuses on primary results from recent decades, and on problems of polygenesis of natural diamond. Correspondingly, we are unable to provide all references addressing this subject, and survey only the most significant publications.

**DIAMONDS IN KIMBERLITES AND LAMPROITES**

*Abundance*

Diamonds from kimberlites and lamproites are the best studied. Among 2500 known kimberlite pipes, only 8–10% pipes are diamondiferous, and only 25 pipes host economic diamond deposits. Total diamond production worldwide from kimberlites and lamproites is 140–150 million carats per year. At the beginning of the 20th century, it was approximately one million carats. Kimberlites occur on all known ancient cratons, except for the North African and Ara-
bian cratons, which, however, harbor diamond placers, suggesting the possible presence of kimberlites (Fig. 1). According to the Clifford’s rule, diamondiferous kimberlites are restricted to ancient Archean cratons, archons (Janse, 1994).

Sources of Diamond in Kimberlites and Lamproites

Diamonds in compositionally different inclusions within kimberlites and lamproites, and paragenetically formed mineral inclusions in diamonds indicate that the primary sources of diamonds are mantle peridotites and eclogites (Sobolev, 1974; Meyer, 1987), and, to a lesser extent, pyroxenites and websterites, from which diamonds were released through partial melting and involved in kimberlite and lamproite magma. Calculated equilibrium conditions indicate that these diamonds were formed at 900—1400°C, at depths of 150—230 km (Boyd and Finnerty, 1980; Stachel and Harris, 2008; Stachel and Luth, 2015; Stachel, 2014; Nimis et al., 2020). Peridotitic diamonds are formed directly in mantle, whereas eclogitic diamonds are generated both through crystallization of mafic magmas at large depths and during subduction of lithospheric plates to depths within the diamond stability field. As a result, diamonds from different sources have different carbon isotope composition ($\delta^{13}$CVPDB).

Carbon Sources. Carbon Isotope Composition of Diamonds

According to the generally accepted concepts, most diamonds’ carbon sources are juvenile carbon-bearing C—O—H fluids. This is corroborated by inclusions of high-density carbonate-bearing fluids (Weiss et al., 2014), carbonated minerals and melts (Kaminsky et al., 2016; Logvinova et al., 2019) in diamonds. To a lesser extent, diamond carbon sources include organic compounds from subsiding slabs and metallic media in the lower mantle.

The carbon isotope composition of natural diamonds is the main tool for deciphering their genesis. E.M. Galimov made an important contribution to this question. He determined for the first time the wide variations of carbon isotope composition up to $-32\%$ $\delta^{13}$CVPDB in diamonds (Galimov, 1984), and in cooperation with N.V. Sobolev et al., he established differing isotope characteristics of peridotitic and eclogitic diamonds (Sobolev et al., 1979). The carbon isotope variations in diamonds from kimberlites and lamproites measured to date span a wide range from $-41$ to $+5\%$ $\delta^{13}$CVPDB (Cartigny et al., 2014). The majority of values fall in the range from $-8$ to $-2\%$ $\delta^{13}$CVPDB, averaging $-5 \pm 1\%$ $\delta^{13}$CVPDB, which is taken to be the average mantle carbon isotope composition. These values correspond to the isotope composition of peridotitic diamonds. Insignificant variations are caused by carbon isotope fractionation during diamond formation.

According to experimental data, the fractionation of carbon isotopes during diamond crystallization from carbonate fluids is $3.9\%$, being no more than $4.5\%$ in the metal—carbon system (Reutsky et al., 2015).

Eclogitic diamonds display much wider $\delta^{13}$CVPDB variations, spanning the entire range from $+5$ to $-42\%$, with maximum in the region from $-20$ to $-28\%$. This is caused by differencing carbonate compound compositions, which undergo subsidence with lithospheric plates at mantle depths, and crystallize there as diamonds, practically without isotope fractionation.

Over 1% of peridotitic diamonds from South Africa (Deines et al., 1993, 2009, and others; Michael et al., 2010), Arkhangelsk province (Galimov et al., 1994), Brazil (Kaminsky et al., 2001; Bulanova et al., 2010), and Canada (Cartigny et al., 2009) have an anomalous carbon isotope composition ($\delta^{13}$CVPDB = $-12$ to $-25\%$). The cause of this phenomenon remains unclear. Some of these diamonds are likely related to the deep mantle horizons, as is suggested by the presence of iron carbide inclusions. This raises the question of Earth initial carbon isotope composition, given that the $\delta^{13}$CVPDB of chondrite varies from $-14$ to $-22\%$ (Wood et al., 2013), while Mars, Vesta, and Moon meteorites have $\delta^{13}$CVPDB = $-20 \pm 4\%$ (Grady et al., 2004). Thereby, the isotope composition of diamond from carbonaceous chondrites varies from $-28.0$ to $-40.2\%$ $\delta^{13}$CVPDB (Russel et al., 1996).

Mechanism of Diamond Formation in Different Media

The majority of diamonds transferred to kimberlites and lamproites from mantle ultramafic rocks were formed through subsolidus metasomatism with the assistance of juvenile C—O—H fluids. According to studies of inclusions in diamonds, the mantle is characterized by reducing conditions, with $\Delta$log $f_O$, varying from fayalite—magnetite—quartz (FMQ) to iron—wüstite (IW) buffer and locally even lower (especially in the lower mantle at the contact with core) (Kaminsky et al., 2015). Under these conditions, the C—H—O fluids are mainly made up of water (90—99 mol %) with a small admixture of CH$_4$ and CO$_2$ (Shirey et al., 2020). The decompression ascent of the mantle with cooling along a geotherm leads to carbon release and isochronal diamond formation, which does not require oxygen exchange with host ultramafic rocks (Stachel and Luth, 2015). At $\Delta$log $f_O$, (FMQ) = $-1.5$…$-2.4$, $T$ = 1140°C, and $P$ = 5 GPa, the following reaction proceeds:

$$\mathrm{CO}_2 + \mathrm{CH}_4 \rightarrow 2\mathrm{C} + 2\mathrm{H}_2\mathrm{O}. \quad (1)$$
Under more reducing conditions \((\Delta \log f_0_2 \text{ (FMQ)} \leq -3)\), diamond can be crystallized from ascending fluids through the following redox neutral reaction:

\[
2\text{C}_2\text{H}_6 \rightarrow 3\text{CH}_4 + \text{C}. \tag{2}
\]

Diamond formation in carbonate medium under upper and lower mantle conditions (5.5–8.4 GPa) has been confirmed by experimental methods (Palyanov et al., 2016; Solopova, 2014).

Some kimberlitic diamonds, including the largest ones, contain metallic inclusions of Fe–Ni–C–S composition, indicating their possible crystallization from a metallic melt with dissolved carbon (Smith et al., 2016a). In addition to the metallic inclusions, the diamonds also contain davemaoite (former CaSi-perovskite), which may indicate their formation in deep Earth. Diamond-forming metallic melts could exist as domains and zones in the lower mantle, especially in the transitional zone D″ at the core–mantle boundary. It is noteworthy that in addition to metals, inclusions may be represented by cohenite \(\text{Fe}_3\text{C}\), which indicates a local carbon oversaturation of the system and could facilitate diamond crystallization.

Data on the carbon isotope composition in these diamonds are interesting. In addition to values of \(\delta^{13}\text{C} = -4\) to \(-9\%\) typical of mantle diamonds, some crystals have a light carbon isotope composition of \(-25.1\) and \(-26.9\%\) (Smith et al., 2016a). It has been hypothesized that this could be evidence of the involvement of crustal carbon in the formation of these diamonds. However, it is possible that such \(\delta^{13}\text{C}\) values could be typical of core carbon.

The mechanism of diamond crystallization from Fe–Ni–C–S melt has been experimentally implemented in the absence (Zhimulev et al., 2012) and presence of a silicate component (Chepurov et al., 2020). Therefore, in diamonds synthesized from metallic melts, metallic inclusions are surrounded by \(\text{CH}_4\) and \(\text{H}_2\) fluids (Smith et al., 2016b), as observed in natural diamonds.

Some deposits, in addition to common diamond crystals, contain small (0.25–0.5 mm) octahedral crystals with flat smoothed faces, usually lacking dissolution traces. These crystals are characterized by a low total nitrogen content (compared to other larger diamonds), with a predominance of \(A\)-defects (nitrogen atomic pairs). The fraction of aggregated nitrogen (%N\(_\text{A}\)) in them is low, near 20%. Such diamonds were found in the kimberlite pipes of the Arkhangelsk province (Palazhchenko, 2008) and in the Madhavan lamproite pipe, India (our observations). We suggest that small octahedral diamond showing no oxidation traces even on the crystal faces were formed at the late stage of kimberlite evolution, already in magma.

Mechanism of diamond crystallization in silicate and silicate–carbonate medium has been experimentally confirmed and studied in detail (Litvin, 2009, and references therein).

**Depth of Diamond Formation in Kimberlites**

Near 90% of diamonds are formed in the subcratonic lithospheric mantle (Stachel and Harris, 2008). Thermobarometric calculations for coexisting mineral inclusions show that they crystallized at depths of 150–230 km, averaging 175–180 km (Stachel, 2014; Nimis et al., 2020), indicating that crystallization occurred in the upper mantle. Thereby, diamonds from different deposits have been formed at practically the same depths, however, temperatures estimates deviate from the geotherm to the higher and lower values.

In the 1990s, majorite, bridgmanite, davemaite (former CaSi-perovskite), ferropericlase, and other minerals were found in diamonds from Brazil (Harte et al., 1999; Kaminsky et al., 2001) and Guinea (Stachel et al., 2000a, 2000b). According to experimental data, these minerals are formed in the transitional mantle zone (220–660 km in depth) and lower mantle (660–2900 km). Reliable geothermometers for these associations are absent, and we can only qualitatively estimate that Fe-rich associations are the deepest ones (Kaminsky and Lin, 2017), while diamonds with metallic inclusions could crystallize in the lowest mantle horizons (Smith et al., 2016a).

**DIAMONDS IN NONKIMBERLITIC VOLCANIC ROCKS**

Diamond finds in lamprophyres, picrites, and analogous volcanic rocks were hotly debatable several tens of years ago, but at present are a common phenomenon. Within the last 15 years, several dozen known occurrences (Kaminsky, 2007) have been supplemented by diamondiferous lamprophyres in the Nunavut (MacRae et al., 1996; Kaminsky and Sablukov, 2002), Northern Ontario (DeStephano et al., 2006; Wyman et al., 2015), and Northern Quebec (Birkett et al., 2004) provinces of Canada; komatitites of French Guiana (Cartigny, 2010; Smith et al., 2016) and Suriname (Naipal et al., 2019), shoshonite–absarokite–picrites of Uzbekistan (Golovko and Kaminisky, 2002), and others occurrences (Fig. 2). Many of them were found within ancient fold belts instead of ancient cratons. Some of them (in French Guiana and Northern Quebec) are subeconomic deposits.

The most economically promising are ultrabasic lamprophyres subdivided into alnoites (withgroundmass melilit), aillikites (with primary carbonate), and damtjernite (withgroundmass nepheline and/or alkali feldspar) (Tappe et al., 2005). All types of the lamprophyres contain diamonds. Mineralogically, they are similar to diamonds from kimberlites and lamproites. As in kimberlites and lamproites, diamonds were formed there under high-temperature and high-pressure mantle conditions. The evidence for such mecha-
nism is the discovery of mantle peridotite xenolith with a diamond inclusion in lamprophyres of North Timan (Kaminsky et al., 1986).

**DIAMONDS IN IMPACTITES**

In the early 1970s, peculiar small (< 1 mm in size) “laminated” diamond crystals were found in the rocks of the 100-km Popigai meteorite crater (Masaitis et al., 1972). Similar diamonds have subsequently been found in other astroblemes in Russia, Ukraine, Germany, Finland, and Canada (Rost et al., 1978; Valter et al., 1992; Masaitis, 2013, and others), as well as in placers of Ukraine (Polkanov et al., 1973) and Yakutia, where they were called “yakutites” (Kaminsky et al., 1985). They were distinguished as independent XI diamond variety (Orlov and Kaminsky, 1981).

Impact diamonds, in addition to their small size, differ sharply from kimberlitic and lamproitic diamonds in their lamellar morphology, inherited from graphite precursor, microcrystalline texture, and, most significantly, an admixture of hexagonal carbon modification, lonsdaleite, amounting from 5 to 70%. The lonsdaleite admixture impart specific properties to diamonds. Owing to the presence of lonsdaleite and nitrogen aggregation, their abrasive properties are 1.5–2 times higher than those of common (natural and synthetic) monocrystalline diamonds (Masaitis, 2013).

Galimov et al. (1980) studied carbon isotope composition of impact diamonds, and established that their composition is identical to the isotope composition of graphite in the target rocks of the Popigai crater, confirming that yakutites were formed through graphite recrystallization.

Diamonds have been found in many of 150 astroblemes. Fifteen structures, in addition to the Popigai crater, have been found in Russia. New types of impact diamonds formed through crystallization of coal, and organic remains have been found in the well-studied ~65 km Kara diamondiferous astrobleme in the Polar Urals (Shumilova, 2021).

Impact diamonds have originated from impact transformation of graphite contained in graphite-bearing gneisses (Popigai) or black shales (Kara). They were formed under dynamic compression at temperatures >1000°C and shock loads of 35–60 GPa for approximately half a second, with subsequent relaxation (Masaitis, 1998). The diamond content in impactites can reach economic concentrations (up to 100 carats/ton) at reserves of tens and hundreds of billions of carats, which is many times higher than the world reserves of common “kimberlitic” diamonds. However, at present the profitability of their mining seems to be ambiguous. The main problem facing the study of impact diamonds is the detailed reconstruction of the conditions under which they form, which
would allow the economic synthesis of such diamonds with their unique abrasive properties.

**DIAMONDS IN METAMORPHIC ROCKS**

In 1970s, diamonds were also found in metamorphic rocks of the Kokchetav Massif, Kazakhstan. Similar diamonds have subsequently been found in many regions worldwide, including in Dabie Shan and Kaidam in China, Norway, Germany, Sulawesi in Indonesia, Greece, South Urals, and other areas (Fig. 3). At the best studied Kumdy-Kol deposit in Kazakhstan, economic diamond concentrations (near 20 carat/ton) have been observed in different lithologies from silicate to essentially carbonate rocks (e.g., Dobrzhinetskaya, 2012). Therefore, metamorphic microdiamonds are frequently present within garnet and zircon grains. As impact diamonds, they are small in size (usually, 100–500 μm) but contain no lonsdaleite admixture. For this reason, metamorphic diamonds lack high abrasive properties and are thus of no practical interest, although reserves of these diamonds, for instance, in the Kokchetav deposit, reach into the billions of carats.

Several hypotheses (mantle, crustal-mantle, and crustal hypotheses) have been proposed to explain the origin of metamorphic diamonds. The generally accepted model is ultrahigh-pressure metamorphism, where crustal rocks with subduction slabs are submerged deep in the mantle (120–150 km and deeper) reaching the diamond stability field, where diamond is crystallized from carbon-bearing sediments, with subsequent exhumation of the metamorphosed diamondiferous rocks (Sobolev et al., 2003; Liou et al., 2007, 2014; Dobrzhinetskaya et al., 2011). Alternative models suggest fluid-assisted metastable diamond crystallization under subsurface conditions (Pechnikov and Kaminsky, 2008).

**DIAMONDS IN OPHIOLITES**

Diamonds were found in ophiolites for the first time in Tibet (Fang and Bai, 1981) and have since been discovered in different regions of the world: in the Inner Mongolia in China, on the Polar Urals in Russia, in Turkey, Armenia, Myanmar, India, Albania, Cuba, and other localities (Fig. 3). In these complexes, diamonds are usually confined to chromitites (Yang et al., 2021).

There is no consensus on the genesis of these diamonds. In particular, the study of minerals in peridotites from Donqiao ophiolites (Tibet, China) has allowed researchers to conclude that diamonds encapsulated in chromite grains were brought up from a depth over 380 km (Xiong et al., 2017). Some scientists have suggested that such high-pressure peridotites, containing chromite, high-pressure minerals, and minerals requiring highly reducing environment can be formed in mid-ocean ridges (Zhou et al., 2014). Other geologists believe that the presence of the rare

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**Fig. 3.** Distribution of diamondiferous metamorphic rocks and ophiolites.
inclusions of zircon, rutile, andapatite in chromitites indicates their origin in subsiding slabs during subduction (Robinson et al., 2015). Some contemporary hypotheses suggest that chromitites are formed at depths of ~5–10 km at temperatures within 1050–1200°C and pressures 1–1.7 GPa (Ridley, 2013).

Galimov proposed a new model for the diamond formation in chromitites. According to this model, ophiolitic nanodiamonds originate in the deep mantle. Then, nuclei are transported in mantle flows to mid-ocean ridges, where they serve as nuclei for crystallization of larger diamonds, which form in chromitites due to interactions between mid-ocean ridge basalt (MORB) melt and host mantle peridotites (Galimov and Kaminsky, 2021). During subsequent spreading and obduction, these diamonds are exhumed with ophiolite complexes.

DIAMONDS IN MODERN VOLCANIC ROCKS

Historical data record the discovery of small diamonds in lavas of modern eruptions in Indonesia (Gunung-Ruang Island), but these reports have not received scientific confirmation. In recent decades, such finds have been noted in Kamchatka. Diamonds were found for the first time in the Icha Volcano in the 1970s (Kutyev and Kutyeva, 1975), and were later discovered in the volcanic rocks of the Avacha, Koryaksky, Klyuchevskoy, and Alaid volcanoes (northern Kuril Range). More recently, diamonds have been found in eruption products of the Tolbachik Volcano (Gordeev et al., 2014; Karpov et al., 2014, and others) (Fig. 2).

The volcanic diamonds of Kamchatka include monocrystals, which resemble diamonds from chromitites in morphology, size, and physical properties, and polycrystalline aggregates (“kamchatites”). Detailed complex studies of Tolbachik diamonds have revealed their natural origin (Galimov et al., 2016, 2020). It has been shown that nitrogen in these diamonds was incorporated in a non-aggregated state, while mineral inclusions contain compositionally unique silicide grains. These diamonds also have elevated concentrations of trace halogens (F, Cl), the distribution of which is similar to that in volcanic gases. This supports the hypothesis that volcanicogenetic diamonds are formed via a process held in common with diamond crystallization at high temperature and pressure in kimberlites (Bundy, 1996), which is most frequently used to characterize the phase composition of carbon.

This diagram shows equilibrium relationships between thermodynamically stable phases: graphite, diamond, and melt, for one-component “carbon” system, i.e., equilibrium boundaries graphite/diamond, graphite/melt, and diamond/melt (solid lines). The position of the main graphite/diamond boundary is determined using the following equation:

\[ P(T) = (2.73T + 2.02 \times 10^3) \text{ MPa} \]  

(3)

However, diamond is not kinetically transformed into graphite under \( P-T \) conditions of graphite field, but enters a thermodynamically metastable phase leading up to the short dashed curve. At higher temperatures, the metastable diamond beyond this boundary is transformed into stable graphite approaching the \( P-T \) conditions of graphite melting. Thus, in nature, the metastable diamond can coexist with stable graphite. The same kinetic process has been observed for metastable graphite in the diamond stability field.

FINDS OF DIAMONDS IN OTHER ROCKS UNDER DIFFERENT CONDITIONS

Diamond finds in nature are not limited by the above-mentioned examples. For instance, microdiamonds have been found in the Akluilak minette dikes (Nunavut territory, Canada) (Galimov and Sablukov, 2002) and in the Precambrian greenstone rocks and dikes of the Wawa subprovince (Northern Ontario, Canada) (De Stefano et al., 2006). These diamonds represent nuclei in the diamondiferous roots of ancient continent (Stachel et al., 2006). Nanodiamonds have been found in pyroxene xenoliths from Hawaiian plume products (Wirth and Rocholl, 2003). They may also be diamond nuclei, formed in the oceanic sublithospheric mantle.

The sources and genesis of peculiar polycrystalline diamonds, carbonado, have remained unknown for over two centuries (Kaminsky, 1994).

These occurrences indicate that the diversity of diamond finds is not exhausted, and that discovery of new surprising new diamond sources should be expected with expansion of prospecting works, scientific studies, and analytical techniques.
Hexagonal symbols show the $P-T$ fields, where lonsdaleite was synthesized by pulsed heating in a static experiment, or by explosion, under the influence of pulsed pressure and temperature. Thus, “hexagonal diamond” (lonsdaleite) is a “kinetic” product of ductile deformations during diamond forming in nature through the impact process. According to thermodynamic criteria, it represents a metastable phase that is kinetically stable to certain extent. Therefore, it has no own field in the thermodynamic equilibrium diagram.

The enthalpy values for transition of 1 mole of carbon atoms from crystalline diamond and graphite structures into a state of gaseous isolated atoms (enthalpy of atomization) are 713 and 715 kJ/mol, respectively. Owing to the fact that the mole energy of a diamond lattice is 2 kJ lower than that of a graphite lattice, diamond can grow in a metastable $P-T$ field together with graphite, especially, in the presence of finest seed serving as a crystallization center. Diamond crystals forming in a metastable field are small in size (nanometers). Therefore, they have only occasionally been found, for instance, in a melt inclusion from garnet peridotite brought to the surface by volcanic products of the Hawaiian plume on Oahu Island (Wirth and Rocholl, 2003).

In the nanosize region, from 3 to 15 nm, a diamond nucleus is a more favorable crystalline form of carbon atoms than graphite. Under disequilibrium conditions, during “quenching”, diamond becomes a metastable phase, but a change of crystal lattice could require billions of years. Thus, diamond retains stability against external mechanical influences. Nanodiamond, graphite, and silica carbide were initially found as relict grains-inclusions in carbonaceous chondrites (Lewis et al., 1989) and were later studied in detail (Lavrukhina, 2004). Their stellar origin is indicated by their extremely anomalous nucleogenic isotope compositions and peculiar inert gases compositions. The nanodiamond concentration in a cosmic space can be determined from Table 1.

This phenomenon can be explained by thermodynamic calculations using a previously developed algorithm for studies of nanoparticles (Jiang and Chen, 2006). Three crystalline carbon polymorphs are well known: diamond (D), graphite (G), and fullerene (F). Carbon atoms have $sp^3$ (tetrahedral) hybridization in diamond and $sp^2$ (trigonal) hybridization in graphite. Therefore, diamond has a 3D structure, whereas graphite consists of two-dimensional carbon layers arranged in an AB stacking sequence. This sequence differs from the AB sequence in the hexagonal close-packed (HCP) structure and is bound by weak Van der Waals $s$-interaction between delocalized $\pi$-orbitals. Under normal conditions and macrosizes ($d$), graphite is a stable carbon polymorph, and $G \rightarrow D$ transition

![Fig. 4. Phase carbon diagram in $P-T$ coordinates. See text for explanation.](image-url)
occurs only at high temperatures and pressures in compliance with the $P$–$T$ equilibrium phase diagram of carbon (Bundy et al., 1994):

As $d$ decreases to nanometers, diamond can be synthesized at lower temperatures and much lower pressure (Sundqvist, 2021). Thereby, it has been noted that diamond, initially formed at $d < 3$ nm, transformed into graphite as it grows, indicating a first-order size-dependent phase transition. In addition, with a further decrease in $d$ to 2 nm, the most stable carbon form could be fullerene, as experimentally observed. This phenomenon has been theoretically simulated with various levels of sophistication, from quantum-mechanical ab initio to thermodynamic calculations (Barnard et al., 2003). Despite the known constraints imposed by thermodynamic considerations, the results show good agreement with experimental data and will be used in our further considerations.

The above-mentioned carbon phase transitions of carbon can be described using the following thermodynamic values: molar surface free energy, $G_S$, caused by the surface energy $\gamma$; molar elastic free energy, $G_E$; caused by surface tension $\ell$; and molar Gibbs free volume energy $G_V$. Because the difference in free energies of phases $i$ and $j$ $\Delta G^{i-j}(T, d)$ for solid particles displays a complex dependence on temperature $T$ and size $d$, it can be presented in simplified form as a sum of separate members: $\Delta G^{i-j}_V(T)$ is the difference of the temperature-dependent molar Gibbs free volume energies; $\Delta G^{i-j}_S(d)$ is the difference of the surface molar free energies; and $\Delta G^{i-j}_E(d)$ is the difference in molar elastic free energies. Each of these parameters depends only on size (Yang and Li, 2008).

Respectively,

$$
\Delta G^{i-j}(T, d) = \Delta G^{i-j}_V(T) + \Delta G^{i-j}_S(d) + \Delta G^{i-j}_E(d). \tag{4}
$$

For the sake of simplicity, we suggest that carbon particle with a molar volume $V_m$ have spherical shape and are isotropic.

The difference of the molar surface free energies of two phases, $\Delta G^{i-j}_S(d)$, can be represented in the following form according to (Zhao et al., 2002):

$$
\Delta G^{i-j}_S(d) = A_m \gamma_S^i d - A_m \gamma_S^j d, \tag{5}
$$

where $A_m$ is the area of the molar phase surface, which can be expressed by the molar volume $V_m$ as $A_m = 6V_m/d$.

The difference of the molar elastic free energy, $\Delta G^{i-j}_E(d)$, can be represented as follows:

$$
\Delta G^{i-j}_E(d) = P_m \frac{V_m^j}{V_m^i} - P_m \frac{V_m^i}{V_m^j}, \tag{6}
$$

and is determined by different surface tensions of the two phases. According to the Young–Laplace equation, $P_m = f/d$, where $f$ for carbon crystalline phases is determined by equation (Jiang et al., 2001)

$$
f = h\left[(S_{vib} H_m)/(2k V_m R)^{1/2}\right], \tag{7}
$$

where $R$ is the gas constant; $H_m$ is the molar enthalpy of fusion; $S_{vib}$ is the vibrational term of the total molar entropy of fusion $S_m$; $k$ is the compressibility; and $h$ is the atomic diameter. Numerical simulation for fullerene C$_60$ using this equation yielded $f = 2.36–4.02$ J/m$^2$. Formula (7) yields $f = 3.54$ J/m$^2$ (Table 2), consistent with this interval.

At equilibrium between the two phases, $\Delta G^{i-j}(T, d) = 0$, and the corresponding critical size of carbon particle $d_{c}^{i-j}(T)$ is determined from equation (4) as follows:

$$
d_{c}^{i-j}(T) = 4fV_m^j - fV_m^i
+ 6(\gamma V_m^i - \gamma V_m^j)\sqrt{G^{i-j}_V(T)}. \tag{8}
$$

The value $\Delta G^{i-j}_V(T)$ can be determined separately for different pairs of carbon phases: diamond–graphite (D–G), diamond–fullerene (D–F), and graphite–fullerene (G–F). In particular, $\Delta G^{i-j}_V(T)$ for the phase

| Phase          | Content, ppm | Size, μm | Isotope anomalies |
|----------------|--------------|----------|-------------------|
| Graphite (G)   | <1           | 1–6      | C, N, O, Mg(Al), Si |
| Diamond (D)    | -400         | ~0.002   | N, Sr, Ba, inert gases |
| SiC            | -5           | -0.2–10  | C, N, Mg(Al), Si, Ca |

| Phase          | Content, ppm | Size, μm | Isotope anomalies |
|----------------|--------------|----------|-------------------|
| Graphite (G)   | <1           | 1–6      | C, N, O, Mg(Al), Si |
| Diamond (D)    | -400         | ~0.002   | N, Sr, Ba, inert gases |
| SiC            | -5           | -0.2–10  | C, N, Mg(Al), Si, Ca |

| Value          | Diamond (D) | Graphite (G) | Fullerene (F) |
|----------------|-------------|--------------|---------------|
| $h$, nm        | 0.154       | 0.142        | 0.157         |
| $V_m$, cm$^3$/mole | 3.417       | 5.398        | 2.7$d$       |
| $H_m$, kJ/mole    | 125         | 120          | 53            |
| $T_m$, K         | 3723        | 4800         | 1600          |
| $S_m$, J/mole K   | 33.58       | 25.00        | 33.12         |
| $S_{vib}$, J/mole K | 6.37       | 4.75         | 6.29          |
| $k$, $10^{-10}$ 1/Pa | 0.088       | 1.000        | 0.212         |
| $E_v$, kJ/mole    | –           | -714         | -675          |
| $f$, J/m$^2$     | 6.10        | 1.10         | 2/98$d^{1/2}$ |
| $\gamma$, J/m$^2$ | 3.70        | 3.27         | 0             |

Table 2. Thermodynamic parameters of carbon nanoforms (after Zhao et al., 2002)
forms depending on their size and temperature. A decrease in the energetic barrier to diamond nucleation is mainly caused by changes in surface tension and related surface energy. The $G \rightarrow D$ transition is driven by inner pressure, caused by surface tension ($f$), which leads to formation of the high-pressure phase, diamond, from nano-sized carbon particle. In turn, the diamond-to-fullerene transition $D \rightarrow F$ can be explained by a decrease in surface energy ($f$). With decreasing size ($d$), carbon clusters are transformed in the following sequence $G \rightarrow D \rightarrow F$, which correspond to the sequence of bond hybridization $sp^2 \rightarrow sp^3 \rightarrow sp^4$.

In general, the conclusions of thermodynamic model are consistent with available geological, experimental, and theoretical results.

**CONCLUSIONS**

This work demonstrates the diversity of different rocks and tectonic conditions in which diamond occurs, and indicates that its formation is not limited to high $P$–$T$ conditions in kimberlite pipes, as was believed 50 years ago. During these years, economic diamond potential has been established for lamproites, lamprophyres, impacities, and metamorphic rocks. The practical importance of diamonds discovered in ophiolites and volcanic rocks of modern eruptions remains to be estimated. There are also other, more enigmatic diamonds. New diamond discoveries in other rock types and geotectonic conditions have not been excluded.

The formation of diamonds in non-kimberlitic rock types is not always caused by crystallization under stable high temperature and high pressure conditions. Conditions remain ambiguous for metamorphic, ophiolitic, and volcanic diamonds, whose formation is likely related to non-traditional mechanisms. Some geological data indicate the possibility of diamond crystallization outside the $P$–$T$ diamond stability field.

Thermodynamic considerations and calculations have confirmed this possibility. One of the decisive factors supporting “metastable” diamond crystallization is the size of formed crystallites. The main roles in decreasing the energetic barrier of nucleation for nano-sized diamonds are played by surface tension and related surface energy. As a result, the graphite-to-diamond transition is mainly controlled by the internal pressure caused by surface tension, which leads to the formation of high-pressure phase, diamond. These diamond nanocrystals could serve as “seeds”, nucleating crystallization of larger diamonds.

Thus, geological and experimental data, confirmed by thermodynamic considerations, indicate that diamond is a polygenetic mineral formed under different physicochemical conditions in different geotectonic settings. Deciphering these conditions is the main task of modern studies, the results of which may be used in the following sequence $G \rightarrow D \rightarrow F$.
searching for new diamond deposits and improving techniques for diamond synthesis.

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