Article

Microrelief of Rounded Diamond Crystals as an Indicator of the Redox Conditions of Their Resorption in a Kimberlite Melt

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Abstract: We conducted a detailed study of the morphology of diamond crystals partially dissolved in a water-bearing kimberlite melt at pressure of 6.3 GPa, temperature of 1400 °C, and two oxygen fugacities (fO2) corresponding to the Re-ReO2 buffer and near the magnetite–hematite (MH) buffer. The triangular etch pits on the {111} faces, which formed during experimental diamond dissolution, were found to completely correspond to negative trigons on natural diamond crystals in the shape and sidewalls inclination angle. Furthermore, two experimental fO2 values were associated with two relief types of the rounded tetrahexahedroid surfaces typical of natural rounded diamonds. Therefore, the surface microrelief on rounded natural diamond crystals was concluded to be an indicator of the redox conditions of natural diamond resorption.

Keywords: diamond; resorption; crystal morphology; kimberlite melt; high-pressure; high-temperature

1. Introduction

At present, no one doubts that the round shape and most surface patterns of natural diamond crystals were formed during the natural resorption process. Natural diamond resorption is supposed to occur in kimberlite magma during diamond transportation from the mantle to the Earth’s surface [1–4]. To date, there is no universal primary kimberlite melt model. The occurrence of silicate–carbonate melt inclusions trapped in megacrysts at different stages of kimberlite magma evolution, confirm that kimberlites magmas are carbonated melts [5]. Most researchers believe that the characteristic features of the composition of primary kimberlite magma are high MgO concentrations, but relatively small amounts of SiO2 (25–35 wt%), Al2O3, and alkalis [6–8]. Kamenetsky et al. [9,10] believes that primary magma was almost anhydrous rich alkalis and halogens carbonate melt. Despite the contradictions, there is consensus in the literature that unique the unserpentinized units of the Udachnaya-East kimberlite may represent an example of “pristinely preserved” kimberlite. Thus, the Udachnaya-East kimberlite can act as an adequate model of primary kimberlite melt. It should be noted that kimberlite melts can be not only a medium for the resorption of diamond, but also for its crystallization [11,12]. Based on experimental data, the conditions of natural resorption of diamond were reconstructed, under which rounded forms of diamond crystals are formed. These forms were shown to emerge upon dissolution of plane-faced diamond crystals in water-bearing carbonate, silicate, and carbonate-silicate melts, or fluids in a wide range of temperatures (1100–1500 °C) and pressures (1.0–6.3 GPa) [13–20]. However, despite similar macromorphology of natural rounded diamonds, their surfaces have different reliefs. The relief is determined by the presence of trigons, serrate and shield-shaped laminae, shagreen,
drop-like hillocks, and many other patterns. Only, for some of these elements of the rounded diamond microrelief, the formation conditions were determined. A number of surface relief elements are unambiguously associated with the features of the internal diamond structure. It has been found that trigons on the [111] faces are located at the dislocation outputs [21–24], and the intersecting striation is associated with plastic deformation bands [3]. Drop-like hillocks forming shagreen or block-type relief of rounded surfaces occur at sites of strong lattice deformation in diamond crystals [25]. Some microrelief elements are associated with conditions of natural diamond resorption. Orientation of trigons at high pressures was demonstrated to be controlled by two oxygen fugacities (fO₂) and process temperature [26] as well as CO₂ (or CO₂²⁺) content in resorption medium [18,27,28]. Comparison of diamonds resorbed at pressures of 1–3 GPa and 5–6 GPa in media of similar composition did not reveal any fundamental differences between them. In both cases, glossy, low relief surfaces with negatively oriented trigons and ditrigonal layers on the [111] faces were formed [14,15,18,29].

In the previous experimental study on simulation of primary kimberlite melt composition, we investigated the specificity of diamond resorption by these melts [19]. Different oxygen fugacity in the melts was shown to correlate with a difference in the morphology of diamond dissolution. In general, diamond dissolution was accompanied by the formation of shield-shaped laminae, trigons on the [111] faces, and rounded tetrahexahedral surfaces at the octahedron edges. However, the tetrahexahedral surfaces were significantly different. They had fine striation in the case of fO₂ controlled by Re-ReO₂ buffer, but had rounded elements at more oxidized conditions in the case of fO₂ between Re-ReO₂ and Fe₃O₄·Fe₂O₃ (MH) buffers. However, Sokol et al. [19] did not study diamond resorption forms in detail. Therefore, the presented work provides a detailed study of the microrelief of crystals produced in the Ud4, Ud5, and Ud6 experiments [19]. The greatest attention has been paid to the microrelief of rounded tetrahexahedral surfaces and [111] faces, namely, to the striation pattern, shape of layers, and features of trigonal etch pits.

2. Materials and Methods

We studied nine diamond crystals produced earlier in experiments with a water-bearing kimberlite melt [19]. The characteristics of starting materials, design of the high-pressure cell, and experimental procedure were in detail described earlier [19]. Here, we only report the main parameters and experimental conditions. Experiments were performed using a multi-anvil, split-sphere apparatus at a pressure of 6.3 GPa, a temperature of 1400 °C, and a run time of 10 h. The mixture consisted of powder of fresh group I kimberlite from the Udachnaya-East pipe in Yakutia and distilled water added with a micro-syringe before the final ampoule assembly. Before the experiments, kimberlite was cleaned from xenogenic material and had the following composition (wt %): SiO₂–27.52, TiO₂–1.35, Cr₂O₃–0.12, Al₂O₃–2.93, Fe₂O₃–3.10, FeO–5.04, MnO–0.15, MgO–27.68, NiO–0.21, CaO–13.32, Na₂O–2.56, K₂O–1.84, P₂O₅–0.41, H₂O–2.49, CO₂–9.43, S–0.33, F–0.14, and Cl–1.32; total–99.94. Three octahedral crystals of natural diamond from kimberlite pipes Udachnaya and Aikhal (Yakutia) were used in each experiment. The crystals were colorless, without inclusions and cracks visible under an optical microscope, with flat [111] faces, and without signs of natural resorption.

In the experiments, double-capsule assembly were used to control oxygen volatility: An internal Pt ampoule containing sample was enclosed in an external container made of a NiO + CsCl mixture (without addition of water). The Pt capsules were lined with Re foil in order to prevent Fe loss and to fix fO₂ in the sample. External NiO + CsCl containers, due to their low fH₂, ensured hydrogen outflow from Pt ampoules with water-bearing samples, which led to their oxidation to a level controlled by the Re-ReO₂ equilibrium. Therefore, oxygen fugacity upon partial oxidation of Re foil (run Ud-6) corresponded to the Re-ReO₂ buffer, which is about one to two log units above the NNO buffer [19]. In the second case, platinum ampoules without Re foil were used (runs Ud-4 and Ud-5). The loss of iron from the kimberlite melt due to its diffusion into platinum amounted to 19%–28% in these experiments [19]. As shown by Brey et al. [30], significant efflux of Fe to the Pt ampoule was accompanied by oxidation of samples, up to fO₂ of the magnetite–hematite (MH) buffer. Given these
data, it may be concluded that diamond dissolution in experiments with Pt ampoules without Re foil (runs Ud-4 and Ud-5) occurred under more oxidizing conditions in the fO$_2$ interval between Re-ReO$_2$ and MH buffers. This was evidenced by the fact that diamond resorption rates in Pt ampoules without Re foil was 1.2–1.4-fold higher than that in ampoules with Re foil. The results of experiments on diamond dissolution are shown in Table 1.

Table 1. Conditions and results of experiments on the dissolution of diamond in the kimberlite melt at 6.3 GPa, 1400 °C, and duration of 10 h.

| Run No | H$_2$O, wt % | fO$_2$       | Diamonds |
|--------|--------------|--------------|-----------|
|        |              | Initial Weight, mg | Weight Loss, mg | Dissolution Rate, mg/h |
| Ud-4   | 10           | Re-ReO$_2$ < fO$_2$ < MH | 1.02 | 0.20 | 0.02 |
| Ud-5   | 12           | Re-ReO$_2$ < fO$_2$ < MH | 1.02 | 0.24 | 0.024 |
| Ud-6   | 12           | Re-ReO$_2$    | 1.12 | 0.17 | 0.017 |

Before the study, diamond crystals were first treated with a hot mixture of nitric and hydrochloric acids (1:3 volume ratio). Further, to remove possible organic contaminants, diamonds were treated in a hot mixture of concentrated H$_2$SO$_4$ and a 30% K$_2$Cr$_2$O$_7$ aqueous solution (3:2 volume ratio). Diamonds were washed by boiling in distilled water after each acid treatment.

Diamond crystals were studied by optical microscopy using a Carl Zeiss Axio Imager Z2m optical microscope (Carl Zeiss Microscopy, Jena, Germany). The differential-interference contrast (DIC) method was used to increase the contrast of the image of the faces relief.

The total interference contrast (TIC) method was used to generate cross-section profiles of shallow etch pits. Deep etch pits were measured using a MII-4 double-reflecting interference microscope (LOMO, Saint Petersburg, Russia). The error in determining the inclination angles of deep pits was estimated at ±5°. Scanning electron microscopy (SEM) of crystals was performed on a Tescan MIRA3 LMU microscope ((Tescan, Brno, Czech Republic) at the Analytical Center for the Multi-Elemental and Isotope Research of the Siberian Branch of the Russian Academy of Sciences.

3. Results

After the experiments, loss of diamond weight ranged from 15% (Ud-6) to 24% (Ud-5). Crystals significantly changed their initial octahedral shape. Rounded stepped surfaces formed at the octahedron edges. The initial {111} faces were significantly reduced and sculptured by etch pits and shield-shaped or ditrigonal laminae. Large and deep etch channels formed on most crystals. Figure 1 shows SEM photographs of typical diamond crystals after the experiments. Even a quick look at the crystals reveals significant differences in the morphology of diamond dissolution forms produced in a water-bearing kimberlite melt under conditions controlled by Re-ReO$_2$ buffer (Figure 1a,b) and under more oxidizing conditions (Figure 1c,d). These differences are manifested primarily in different sculptures on newly formed rounded surfaces and in a different number of etch pits.

We focused our study on these two relief types of diamond crystals. All faces of each diamond crystal were examined under an optical microscope using the DIC method. Similarly, all crystals were tested in different orientations using a scanning electron microscope (Figure 1). The most representative {111} crystal faces and stepped surfaces were investigated by the TIC method using an optical microscope. Sizes of all detected etch pits were determined, and 40 measurements of the etch pit sidewalls inclination angle relative to the [111] (α[111]) faces were made.
Figure 1. SEM micrographs of diamond crystals after dissolution runs in kimberlite melt at different two oxygen fugacities (fO$_2$): (a,b) run Ud-6; (c) run Ud-4; and (d) run Ud-5.

3.1. {111} Faces

The main relief elements of {111} faces are triangular etch pits and ditrigonal (or shield-shaped) layers. Large layers outlining the {111} faces have an almost triangular shape (Figures 1 and 2). The obtuse angles of the ditrigonal layers are directed to the octahedron edges and have values of approximately 175°. The acute angles of these layers are directed to the octahedron vertexes and have values of approximately 65°. The shape of the layers change toward the center of the {111} faces. The layers have an almost hexagonal shape in the central parts of {111} faces (Figure 2c). Near the <100> vertices, the polycentric development of the layers led to the formation of serrate surfaces.

Figure 2. Optical micrographs (differential-interference contrast (DIC) method) of {111} faces of diamond crystals: (a–c) run Ud-5; (d–f) run Ud-6.
Etch pits on the \{111\} faces have a triangular contour that is oriented 180° relative to the \{111\} faces (Figure 2). In accordance with the terminology used to describe the morphology of natural diamond, we call these etch pits as negative trigons \[31\]. Two types of trigons are found on the studied crystals. Trigons of the first type have a non-truncated pyramidal shape and are called pyramidal and point-bottomed trigons, or, short, p.b.trigons. Another trigon type is a truncated triangular pyramid, i.e., it has a flat base (bottom) parallel to the \{111\} face. They are called flat-bottomed trigons, or, in short, f.b.trigons. We will use this terminology in the rest of the paper.

Figure 2 shows the most representative images of trigons found on the \{111\} faces. Most trigons are symmetrical and from a few to 100 \(\mu m\) in size. The walls of large trigons are usually stepped to varying degrees and, therefore, have a variable inclination to the \{111\} face in most cases. Visual examination under an optical microscope reveals lighter trigons with sloping walls. These trigons include all p.b.trigons and small f.b.trigons. Most large f.b.trigons have dark sidewalls, which indicates their steeper inclinations to the \{111\} face.

Trigonal sidewall inclination angles \((\alpha_{\{111\}})\) were determined using the TIC method. Figure 3 shows typical trigon profiles. The inclination angles of shallow p.b.trigons and f.b.trigons range from 0.5° to 3.5°. Steep non-stepped (or thin-stepped) f.b.trigons have \(\alpha_{\{111\}}\) of about 10°. No differences were found in the angles of \(\alpha_{\{111\}}\) trigons on diamond crystals dissolved at various fO\(_2\) values.

The histograms shown in Figure 4 show the size distribution of p.b.trigons and f.b.trigons on the \{111\} faces of six diamond crystals. Three crystals are partially dissolved at fO\(_2\) of the Re-ReO\(_2\) buffer (run Ud-6), and three crystals are partially dissolved under more oxidized conditions, near the MH buffer (run Ud-5). On diamonds from run U-6, the predominant type is p.b.trigons accounting for 87% of the total trigons. Their maximum size reaches 50 \(\mu m\), but prevailing p.b.trigons (96%) are less than 15 \(\mu m\) in size (Figure 4c). Flat-bottomed trigons are less numerous. Mainly, these are not deep trigons, no larger than 35 \(\mu m\) in size (Figure 4d). On diamonds from run Ud-5, the number of both trigon types is approximately the same and is 55% for p.b.trigons and 45% for f.b.trigons. Unlike run Ud-6, pyramidal trigons are smaller. Most of them (96%) are less than 10 \(\mu m\) in size (Figure 4a). In addition, unlike run Ud-6, the dimensions of f.b.trigons demonstrate a distinctive bimodal distribution.
indicating the presence of two f.b.trigon types (Figure 4b). Relatively small f.b.trigons (with a maximum at 5 µm) have shallow walls. Large f.b.trigons, ranging from 30 to 100 µm in size and with a maximum of 60–70 µm, have steeper walls.

![Figure 4](image)

**Figure 4.** Histograms of the sizes of trigons on the (111) faces of diamond crystals dissolved under various conditions: (a,b) under oxidizing conditions in the fO2 interval between Re-ReO2 and magnetite–hematite (MH) buffers (runs Ud-4 and Ud-5); (c,d) under conditions of the Re-ReO2 buffer (run Ud-6).

3.2. Rounded Surfaces at Octahedron Edges

As already noted, curved rough-relief surfaces formed in place of the octahedron edges. They are characterized by a significant curvature in the direction of {111} faces and an insignificant curvature in the direction of <100> vertices (Figure 1). In the central part of the surfaces, there is a break in the form of a curved line (pseudo-edge) connecting the adjacent {111} faces (Figure 5c,d). These pseudo-edges are most clearly seen on the rounded surfaces of diamonds from Ud-4 and Ud-5 experiments (Figure 5c,d).

Among natural rounded diamonds, these crystals are called tetrahexahedroids or, more often, dodecahedroids. Depending on the redox conditions used in our experiments, the structure of these curved surfaces is significantly different. Diamond dissolution under redox conditions controlled by Re-ReO2 buffer (run 6) led to the formation of rounded stepped surfaces with relatively uniform parallel striation (Figure 5a,b). Near the <100> vertices, parallel striation turns into serrate striation. A TIC study of the surfaces showed that the height of most steps varies from 20 to 150 nm, with a mean of 50–60 nm (Figure 6a–c). Single macrosteps have a height of up to 5–6 µm.

Upon more intensive diamond dissolution under more oxidizing conditions (runs Ud-4 and Ud-5), surfaces with rounded step relief were formed at the octahedron edges (Figure 5c,d). The structure of these surfaces can be characterized as an alternation of rough-stepped relief with relatively smooth rounded areas and very elongated hillocks. Rounded, relatively smooth areas of the surfaces have fine striation; the height of striation steps is not more than 20 nm (Figure 6d–f), with a mean height being several nanometers. Macrosteps and elongated hillocks are up to 10 µm thick.
Figure 5. SEM micrographs of curved rough-relief surfaces formed in place of the octahedron edges: (a,b) run Ud-6; (c) run Ud-4; and (d) run Ud-5.

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4. Discussion

4.1. Trigons on Natural Diamonds

Negatively oriented triangular pits, the so-called negative trigons, are almost always present on the {111} faces of natural diamonds. The walls of trigons are usually stepped and they correspond to a series of trisoctahedra faces. According to the data of a goniometric study of natural diamond crystals from kimberlite pipes of Yakutia, the structure of trigons can involve almost all possible trigon-trioctahedra, with the most common faces being {998}, {775}, {221}, {441}, and {661} [32]. Based on goniometric measurements of trigons on diamonds from the Ural placers, A.A. Kukharenko [33] reported a slightly
different set of the most common trisoctahedron faces. These are mainly trisoctahedrons: {887}, {776}, {665}, {554}, and {443}. However, given light diffraction inevitably occurring upon light reflection from thin-stepped surfaces, simple forms observed using optical goniometry cannot be considered really existing. More reliable information about the trigon structure is obtained using interference methods. With the use of multibeam interferometry has been shown that the walls of shallow p.b.trigons form an α(111) angle of 5° to 2.5° with the {111} face [23,31]. Sometimes, step trigons, whose walls correspond to the {332} faces (α(111) = 10°), are observed [34]. Steep-wall non-stepped f.b.trigons are formed by the {111} faces only [35].

The structure of trigons on natural diamonds can be studied in more detail using atomic force microscopy (AFM). Y. Fedorchouk [28] presented data on the study of trigons on three crystals of diamond from Ekati Diamond Mine kimberlites using AFM. It was found that p.b.trigons from 3 to 40 µm in size have α(111) from 3° to 5°, which can correspond to simple Miller indices—{998}, {887}, {776}, and {665}. According to AFM measurements, walls of f.b.trigons of up to 40 µm in size have α(111) = 7°, 9°, 17°, 19°, and 23° [28]. Given the measurement accuracy, these inclination angles may correspond to {443}, {332}, {944}, {552}, and {331} faces. Comparison of the results obtained by different measurement methods demonstrates good agreement for the inclination angle of negative trigon walls.

Thus, several types of negative trigons are formed on the {111} faces of diamond crystals during natural resorption: (1) shallow trigons, primarily point-bottomed, with sidewalls having α(111) from 5° to 5°; (2) flat-bottomed stepped trigons confined by walls with α(111) from 7° to 23°; and (3) flat-bottomed non-stepped trigons composed exclusively of {111} faces. An unambiguous relationship between p.b.trigons and dislocations was revealed using X-ray topography [21–23] and photoelasticity [24] techniques. All dislocations associated with trigons were found to be rectilinear and full edge dislocations or dislocations with a significant edge component [24].

It is considered unambiguously proven that trigons are formed during natural dissolution of diamond, and, therefore, they are etch pits. The inclination of etch pit walls is determined by the ratio of normal and tangential dissolution rates: \( \tan \alpha = N/V_t \), where \( N \) is the dissolution rate component perpendicular to the surface, i.e., dissolution along the dislocation line, and \( V_t \) is the rate of dissolution parallel to the surface. \( V_t \) is controlled by the rate of step propagation along the surface. This issue is most thoroughly addressed in [36–38]. The application of theoretical models to trigonal pits on diamond was recently demonstrated in [20]. The rate ratio and, therefore, the inclination of etch pit walls depend on many parameters. For example, it was experimentally demonstrated that the inclination of pits walls on diamond may depend both on external parameters (medium composition, temperature, etching time, and oxygen fugacity) and on the features of diamond, such as the type of defects on which etch pits form [39,40] and content of impurities in diamond crystals [41,42]. Under constant etching conditions (medium composition, temperature, etc.), the tangential dissolution rate, \( V_t \), is constant. In this case, the inclination of etch pit sidewalls is mainly controlled by the dissolution rate along the dislocation line (\( N \)). If etching conditions are changed, the tangential etch rate also changes. For this reason, etch pits at the same dislocations have different inclinations. In this respect, of great interest are studies of diamond etching. For example, etch pits on [100] diamond faces, which are formed with participation of oxygen or water vapor, have a wall inclination angle of 3–7° [43,44]. Upon diamond etching with predominantly hydrogen plasma, etch pits of the same shape are formed on the [100] faces, but the sidewall inclination angle amounts to 21–31° [45]. In addition, the shape and inclination of etch pit sidewalls were demonstrated to depend on oxygen fugacity [17,26]. Therefore, the inclination of etch pit sidewalls may be an indicator of etching conditions. In this regard, the experimental reproduction of etching pits similar to natural trigons may indicate the similarity of the experimental parameters with the conditions of resorption of natural diamond.

The features of trigonal etch pits have received little attention in most experimental studies of diamond etching. Usually, researchers confine themselves to describing the shape and orientation of trigonal pits. Table 2 shows the data from studies on diamond dissolution at high temperatures and pressures, in which have investigated trigons in detail. The presented data indicate that etch pits on
the [111] faces, which were produced by diamond dissolution in water-bearing carbonate and silicate melts in previous works [13,16,20] and in the present work, most closely correspond to trigons on natural diamonds. This suggests that water-bearing carbonate and silicate melts and PT-conditions quite accurately reproduce natural diamond resorption at mantle conditions and during diamond transportation to the Earth’s surface.

### 4.2. Tetrahexahedroid Surfaces on Natural Diamonds

The tetrahexahedroid surfaces on natural diamonds have been unambiguously revealed to form during natural resorption. Apparently, depending on the resorption conditions and internal structure features, the rounded surfaces of natural diamonds have different sculptures. The most common sculpture types are as follows: Macroscopically smooth, glossy surfaces; stepped relief with various intensities of sheaf-like striation, from thin-stepped relief to rounded-stepped relief. Clusters of hillocks form block-type and drop-like surfaces, serrate relief, and shagreen microrelief.

In our experiments, we reproduced two relief types of the rounded tetrahexahedroid surfaces. Upon diamond dissolution in a water-bearing kimberlite melt at $P = 6.3$ GPa and $T = 1400$ °C and oxygen fugacity in the range from Re-ReO$_2$ to MH buffer does not significantly affect the trigonal sidewall inclination. However, when diamond dissolves under conditions of oxygen fugacity near the MH buffer, a larger number of large flat-bottom trigons are formed.

#### Table 2. Experimental conditions of runs and characteristics of experimentally obtained and natural trigons on the [111] faces of diamond crystals.

| Refer. | Composition of the Etchant | $P$, GPa | $T$, °C | $\alpha_{(111)}$ f.b.Trigons | $\alpha_{(111)}$ p.b.Trigons |
|--------|---------------------------|----------|--------|-----------------------------|-----------------------------|
| [28]   | CaCO$_3$ + SiO$_2$        | 1        | 1150–1350 | Pt. 6°, 10°, 11°, 15°, 19° | Pt. 6°, 8°, 10°, 12°        |
| [28]   | Mg(OH)$_2$                | 1        | 1150–1350 | 10–11°, 15–16°, 19–23°, 43° | 11°, 19°                    |
| [20]   | H$_2$O $+$ MgO $+$ SiO$_2$| 1–3      | 1150–1400 | 0.6° to 67°, 0.5° to 43°   |                             |
| [13]   | basalt + H$_2$O lamproite | 2.5–5.5  | 1100–1450 | 1–2°, 6–9°, 11–14°, 15–18°, 19–22° |                             |
| [27]   | CaCO$_3$                  | 7.0      | 1700–1750 | Pt. 5°, 13°                  |                             |
| [18]   | Na$_2$CO$_3$ + CO$_2$     | 5.7      | 1400     | Pt. 11.5°, 19.5°, 23.5°     | Pt. 19.5°, 23.5°            |
| [18]   | CaMg(CO$_3$)$_2$ + CO$_2$ | 7.5      | 1600     | Pt. 11.5°, 19.5°, 23.5°     | Pt. 19.5°, 23.5°            |
| [18]   | Na$_2$CO$_3$ + H$_2$O     | 5.7      | 1400     | 1–2°, 7.5°, 12.5°, 16°, 20.5° |                             |
| [16]   | CaMgSi$_2$O$_3$ + H$_2$O  | 5.7      | 1400     | 7°, 11°, 16°, 21°, 60°32°   | 1°–2°                       |
|        | CaCO$_3$ + H$_2$O         | 5.7      | 1400     | 1°–2.5° and 10°              | 0.5°–3.0°                   |
| Present work | Kimberlite + H$_2$O | 5.7      | 1400     | 7°, 10°, 17°, 19°, 23°, 60°32° | 5°–2.5°                     |
| [23,28,31,35] | Natural resorption of diamond | 5.7      | 1400     | 3°–5°                        |                             |

Pt.—positively oriented trigons.

Our experiments showed that upon diamond dissolution in a water-bearing kimberlite melt at $P = 6.3$ GPa and $T = 1400$ °C and oxygen fugacity in the range from Re-ReO$_2$ to MH buffer does not significantly affect the trigonal sidewall inclination. However, when diamond dissolves under conditions of oxygen fugacity near the MH buffer, a larger number of large flat-bottom trigons are formed.

### 4.2. Tetrahexahedroid Surfaces on Natural Diamonds

The tetrahexahedroid surfaces on natural diamonds have been unambiguously revealed to form during natural resorption. Apparently, depending on the resorption conditions and internal structure features, the rounded surfaces of natural diamonds have different sculptures. The most common sculpture types are as follows: Macroscopically smooth, glossy surfaces; stepped relief with various intensities of sheaf-like striation, from thin-stepped relief to rounded-stepped relief. Clusters of hillocks form block-type and drop-like surfaces, serrate relief, and shagreen microrelief.

In our experiments, we reproduced two relief types of the rounded tetrahexahedroid surfaces. Upon diamond dissolution in a water-bearing kimberlite melt at $fO_2$ of the Re-ReO$_2$ buffer, the octahedron edges are truncated by surfaces with well-defined striation relatively uniformly spread through the entire tetrahexahedroid surface. However, when diamond is dissolved under more oxidizing conditions, the octahedron edges are truncated by roughly sculptured surfaces with alternation of rough-stepped relief, relatively smooth rounded areas, and very elongated hillocks. Possibly, wide variations in the relief of rounded natural diamonds from different deposits are associated with variations in redox conditions occurring during kimberlite magma formation in the Earth upper mantle. According to existing estimates, oxygen fugacity of kimberlite magma ranges from NNO-5 to NNO+6 [46]. This corresponds to oxygen fugacity in the range from approximately IW buffer to MH buffer at pressure of 6.0 GPa and temperature of 1400 °C. The run U-6 was carried out at an oxygen fugacity corresponding to Re-ReO$_2$ buffer, which is approximately one to two log units above the NNO buffer. Runs Ud-4 and Ud-5 occurred under more oxidizing conditions in the $fO_2$ interval...
between Re-ReO$_2$ and MH buffers. These two experimental fO$_2$ values were associated with two relief types of the rounded tetrahexahedroid surfaces typical of natural rounded diamonds. Thus, our data indicate wide variations in the morphology of diamonds upon dissolution at fO$_2$ values typical of kimberlite magma.

5. Conclusions

We studied the morphology of diamond crystals partially dissolved in a water-bearing kimberlite melt at pressure of 6.3 GPa, temperature of 1400 °C, run time of 10 h, and two oxygen fugacity values (fO$_2$) corresponding to the Re-ReO$_2$ buffer and near the HM buffer. The formed triangular etch pits on [111] faces were found to be similar to negative trigons on natural diamond crystals in the shape and wall inclination angles. When diamond is dissolved under more oxidizing conditions, a greater amount of relatively large flat-bottomed trigons form. Furthermore, two experimental fO$_2$ values are associated with two relief types of the rounded tetrahexahedroid surfaces characteristic of natural rounded diamonds. When diamond is dissolved in a water-bearing kimberlite melt at fO$_2$ of the Re-ReO$_2$ buffer, the octahedron edges are truncated by surfaces with well-defined striation through the entire tetrahexahedroid surface. When diamond is dissolved under more oxidizing conditions, the octahedron edges are truncated by rough-sculptured surfaces with alternation of rough-stepped relief, relatively smooth rounded areas, and very elongated hillocks. These findings suggest that the surface microrelief of rounded natural diamonds is an indicator of fO$_2$ of natural diamond resorption.

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