

THE LAW OF DIAMOND CRYSTALLOGENESIS.

Volodymyr Kaplunenko

"Nanomaterials and Nanotechnologies Ltd",

Ukraine

e-mail: kaplunenko@gmail.com

Mykola Kosinov

"Nanomaterials and Nanotechnologies Ltd",

Ukraine

e-mail: nkosinov@ukr.net

***Abstract.** In our work [19], we presented the law of diamond crystallogenesis applicable to the synthesis of natural and laboratory-produced diamonds. Here, we describe the law of diamond crystallogenesis in more detail. The law of diamond crystallogenesis shows that the rate of diamond growth depends on the number of electrons involved in the synthesis and the oxidation state of carbon in the precursor. Pressure and temperature play a supporting role and are triggers that initiate the electronic mechanism of diamond synthesis. A key aspect of the law of diamond crystallogenesis is electrons, which are considered the most important catalysts in the process of diamond formation, playing a fundamental role in changing the reactivity of carbon and forming the diamond structure. The discovery of the law of diamond crystallogenesis dispels the myth of millions and billions of years required for diamond formation, as well as the myth of pressure and temperature as direct factors in diamond formation. The law of diamond crystallogenesis opens the way to breakthrough technologies for the ultra-fast synthesis of artificial diamonds.*

***Keywords:** diamond synthesis, law of diamond crystallogenesis, electron as a catalyst, oxidation degree concept.*

1. Introduction

The first theoretical justification for diamond synthesis was Leipunskii's work [1], which laid the foundation for diamond synthesis, specifying the pressure and temperature conditions under which diamond is stable. Leipunskii's diagram was refined and improved by Bundy F. P. [2, 3].

Leipunskii's thermodynamic calculations were confirmed by practical experiments in the 1950s and 1960s [4]. A method for producing artificial diamonds, HPHT, was developed. This served as a convincing reason to believe that pressure and temperature are essential conditions for diamond synthesis. Moreover, pressure and temperature began to be considered direct factors in diamond formation.

At the same time, an increasing number of studies are describing new methods of diamond synthesis that do not require high pressures and temperatures [5-11]. In [5, 6], it was shown that diamonds can be synthesized from adamantane solely using an electron beam. This new method for synthesizing nanodiamonds involves exposing adamantane (a hydrocarbon $C_{10}H_{16}$) to an electron beam at low temperatures, which breaks the C–H bonds, forming a diamond lattice. Nanodiamond synthesis occurs in a vacuum and lasts a short time—tens of seconds. This breaks all the rules and

traditional understanding of diamond formation and contradicts the predictions given in the Leipunsky and Bundy diagrams. This new method for synthesizing nanodiamonds demonstrates that extreme temperatures and pressures are not required for diamond formation. Only electrons in a vacuum are required to transform adamantane into diamond.

Controversies have also arisen in the concept of natural diamond formation. Increasing geological data indicate the possibility of diamond crystallization outside the P–T stability region of diamond [12]. Diamonds have been discovered in rocks that do not contain high-pressure minerals [13–15]. These discoveries in geology and laboratory synthesis demonstrate the inconsistency of the concept of diamond formation based on extreme pressure and temperature. A paradoxical situation has arisen: on the one hand, we have Leipunsky and Bundy's calculations, confirmed by the HPHT synthesis method, which indicate the need for extreme pressures and temperatures. On the other hand, we have new geological data and new diamond synthesis methods that do not require extreme pressure and temperature and allow diamonds to be produced outside the P-T stability region. This contradiction requires an explanation. To resolve it, we must consider the fact that the HPHT method actually involves pressure and temperature gradients, which are not represented in the Leipunsky and Bundy diagrams.

2. The prevalence of empirical research in artificial diamond synthesis, rather than the scientific method.

Currently, laboratory diamond synthesis is more of an art than a science. The selection of catalysts for diamond synthesis lacks a rigorous scientific basis, and their preparation recipes must describe all steps in detail to ensure reproducibility. This is typical for catalytic processes. A similar situation has long been known in catalysis [16-18]. Diamond synthesis as a catalytic process is plagued by the same problems. A successful "recipe" must be developed through trial and error, rather than based on a scientific analysis of the processes occurring during synthesis. The statement cited in [18] is entirely applicable to laboratory diamond synthesis: *"it is more like the art of cooking than stoichiometric chemical synthesis."*

The development of a scientific theory of diamond formation is a pressing issue. This is driven by recent breakthrough discoveries in diamond synthesis methods, described in [5, 6, 7]. This is driven by the contradictions of the modern concept of diamond formation, based on extreme pressure and temperature. Perhaps we won't have to wait long for a quantitative theory of diamond formation that could universally, using a single equation, describe the process of diamond synthesis from various precursors and indicate the parameters that determine the rate of diamond growth. The lack of a scientific theory of diamond formation hinders the development of diamond synthesis technologies. The lack of a scientific theory of diamond formation has given rise to two myths: the myth of millions and billions of years required for diamond formation, and the myth of pressure and temperature as direct factors in diamond formation.

3. The law of diamond crystallogenesis.

In our work [19], we presented a unified mechanism for diamond crystallogenesis in nature and in laboratory synthesis, in which electrons are the primary factor. The electron mechanism of diamond synthesis demonstrates stable, repeating characteristics between physical quantities and

regularly repeating states of the material objects involved in the synthesis. This indicates the existence of a physical law of crystallogenesis that can be represented in a rigorous mathematical formulation. The existence of a physical law of crystallogenesis is also indicated by the fundamental nature of similar physical processes in different synthesis methods, and by the fundamental particle—the electron—that participates in them. The law of diamond crystallogenesis is as follows:

$$v_D = \frac{e \cdot M_e}{F \cdot t \cdot (q_C - q_{C_{\min}})}$$

Fig. 1. The law of diamond crystallogenesis. v_D is the rate of diamond formation (mol/s); e is the electron charge; M_e is the number of free electrons participating in the synthesis; F is the Faraday constant; q_C is the oxidation state of carbon; $q_{C_{\min}}$ is the minimum oxidation state of carbon (-4); t is the time of diamond synthesis.

The physical law of diamond crystallogenesis follows from the mechanism of diamond crystallogenesis, which is based on the interaction of electrons with atomic carbon and the donor-acceptor interaction of C^{-4} carbon atoms with the diamond surface, forming covalent bonds via the donor-acceptor mechanism.

The law of diamond crystallogenesis demonstrates its universality and applicability to all methods of diamond synthesis. It is represented by a simple and elegant mathematical formula. The law of diamond crystallogenesis includes fundamental physical constants that characterize the participants in diamond synthesis. These participants in diamond synthesis are electrons and carbon atoms.

The main player in the mechanism of diamond formation is the electron. It is represented in the formula of the law of diamond crystallogenesis by a fundamental physical constant: the elementary charge " $e = 1.60217663 \times 10^{-19}$ C." From the law of diamond crystallogenesis it follows that the rate of diamond formation is proportional to the number of M_e electrons involved in the synthesis.

This law states that electrons are the primary driving forces in diamond growth, lowering the oxidation state of carbon to C^{-4} , which facilitates the formation of the diamond crystal structure. This is a revolutionary approach to understanding diamond synthesis in both natural and laboratory conditions. It suggests new technologies for ultra-fast diamond production.

A key aspect of the law of diamond crystallogenesis is that it considers electrons to be the most important catalysts in the diamond formation process, playing a fundamental role in the transformation of carbon into its diamond structure. The mechanism of diamond crystallogenesis involves electrons, which change the oxidation state of carbon from higher positive or neutral states to the C^{-4} state required for diamond synthesis. This electronic transformation of carbon's reactivity "involves" the Coulomb interaction in the synthesis mechanism and initiates the diamond synthesis reaction.

Discoveries arising from this law could lead to new technological approaches that could significantly accelerate the process of growing diamond crystals, potentially reducing laboratory processes to just a few hours.

The law of diamond crystallogenesis (Fig. 1) includes Faraday's constant ($F = 9.64853321233100184 \times 10^4 \text{ C/mol}$). This fundamental constant is a physicochemical constant. It links physical and chemical quantities: the electron charge and Avogadro's constant. Faraday's constant is known to be part of the Nernst equation, the Goldman equation, and Faraday's law of electrolysis. The law of diamond crystallogenesis is a physicochemical law that also includes Faraday's constant. The law of diamond crystallogenesis points to the key role of the oxidation state of carbon in diamond synthesis. The law of diamond crystallogenesis suggests that the highest rate of diamond formation is achieved using carbon-containing substances in which the degree of carbon oxidation is minimal. This is observed in practice when using methane (CH_4) as a diamond-forming gas.

Oxidation states are not considered physical quantities. They are chemical characteristics of substances. Their presence in a physical law is unusual, as oxidation states are typically used in chemistry to describe the reactivity of substances. However, oxidation state is included in the law of diamond crystallization as a parameter, along with fundamental physical constants. This is despite the fact that oxidation state is not considered a fundamental parameter in chemistry.

The history of the oxidation state concept spans approximately 200 years [20–23]. Despite its widespread use in chemistry, oxidation state is considered an auxiliary, conventional quantity with no physical meaning [24–27]. Scientists have long debated the role of oxidation state in chemistry [26, 28–30]. Some authors point to the universality and fundamental nature of oxidation state [31–33]. Pauling famously stated, "*If scientific progress continues, the next generation may have a theory of valence that is sufficiently precise and powerful to allow chemistry to be considered an exact science, on a par with physics*" [31, 32]. We regard oxidation states as a fundamental characteristic of substances that determines their reactivity. The inclusion of carbon oxidation states as parameters in the law of diamond crystallogenesis, along with fundamental physical constants, indicates the fundamental status of oxidation states.

4. Conclusion

The presented law of diamond crystallogenesis demonstrates that electrons and atomic carbon in the minimum oxidation state of C^{-4} participate in diamond formation. Electrons act as catalysts, enabling the synthesis of diamond from precursors. They play a fundamental role in changing the reactivity of carbon and in the formation of the diamond structure. Electrons convert the precursor carbon to the lowest oxidation state of C^{-4} , which is necessary for the diamond synthesis reaction to occur. This law is expressed mathematically. It relates the rate of diamond formation to the number of electrons involved in the synthesis, the Faraday constant, the oxidation state of carbon, and time. The law of diamond crystallogenesis emphasizes the central role of electrons in the diamond synthesis reaction, in particular their participation in the activation of the Coulomb interaction necessary for diamond formation. Diamond formation processes previously considered to require extreme conditions can now be realized via an electron mechanism, paving the way for new, ultra-fast diamond synthesis technologies.

5. Conclusions

1. The law of diamond crystallogenesis states that the rate of diamond growth depends on the number of electrons involved in synthesis and the oxidation state of carbon in the precursor.

2. A key aspect of the law of diamond crystallogenesis is that electrons are considered the most important catalysts in the diamond formation process, playing a fundamental role in changing the reactivity of carbon and in the formation of the diamond structure.

3. Electrons convert carbon to the lower oxidation state C^{-4} , which is necessary for the diamond synthesis reaction to occur.

4. The law of diamond crystallogenesis is expressed mathematically. It relates the rate of diamond formation to the number of electrons involved in synthesis, the Faraday constant, the oxidation state of carbon, and time.

5. Diamond formation processes, previously thought to require extreme conditions, can now be achieved under normal conditions using an electronic mechanism, paving the way for new ultra-fast diamond synthesis technologies.

References.

1. Leipunskii O. I. (1939). Ob iskusstvennykh almazakh [On Synthetic Diamonds]. *Uspekhi khimii — Russian Chemical Reviews*, vol. 8, iss. 10, pp. 1519 – 1534.

2. Bundy, F. P. (1962). "Direct Conversion of Graphite to Diamond in Static Pressure Apparatus". *Science*. 137 (3535): 1057–1058. Bibcode:1962Sci...137.1057B. doi:10.1126/science.137.3535.1057. PMID 17774419.

3. Bundy, F. P. (1963). "Direct Conversion of Graphite to Diamond in Static Pressure Apparatus". *The Journal of Chemical Physics*. 38 (3): 631–643. Bibcode:1963JChPh..38..631B. doi:10.1063/1.1733716

4. Valeriy Sobolev, Svetlana Gubenko, Oleh Khomenko, Maksym Kononenko, Roman Dychkovskiy, Adam Smolinski, Physical and chemical conditions for the diamond formation, *Diamond and Related Materials*, Volume 151, 2025, ISSN 0925-9635, <https://doi.org/10.1016/j.diamond.2024.111792>.
(<https://www.sciencedirect.com/science/article/pii/S0925963524010057>)

5. Fu J, Nakamuro T, Nakamura E. Nanodiamond synthesis from adamantane by electron-beam driven C–H activation at 100 K. *ChemRxiv*. 2025; doi:10.26434/chemrxiv-2024-fk6qk-v2

6. Jiarui Fu, Takayuki Nakamuro, Eiichi Nakamura, "Rapid, low-temperature nanodiamond formation by electron-beam activation of adamantane C–H bonds," *Science*: September 4, 2025, doi:10.1126/science.adw2025.

7. Gong, Y., Luo, D., Choe, M. et al. Growth of diamond in liquid metal at 1 atm pressure. *Nature* 629, 348–354 (2024). <https://doi.org/10.1038/s41586-024-07339-7>

8. Y. G. Gogotsi, K. G. Nickel. Hydrothermal Synthesis of Diamond: Challenges and Opportunities
<https://doi.org/10.1002/9780470294444.ch88>

9. DeVries, R. Diamonds from warm water. *Nature* 385, 485 (1997). <https://doi.org/10.1038/385485a0>

10. Chen, C. & Chen, Q. Recent development in diamond synthesis. *Int. J. Modern Phys. B IJMPB* 22, 309–326. <https://doi.org/10.1142/S0217979208038685> (2008).

11. Andrzej M. Szymanski. On the Possibility of Bulk Large Diamond Single Crystal Synthesis with Hydrothermal Process. *Journal of the Mineralogical Society of Korea*, v.10 no.1, pp.18-32, 1997.
12. Kaminsky, F.V., Voropaev, S.A. Modern Concepts on Diamond Genesis. *Geochem. Int.* 59, 1038–1051 (2021). <https://doi.org/10.1134/S0016702921110033>
13. Farre-de-Pablo J., Proenza J.A., Gonzalez-Jimenez J.M. et al. A shallow origin for diamonds in ophiolitic chromitites // *Geology*. 2018. Vol. 47. № 1. P. 75-78. DOI: 10.1130/G45640.1
14. Pujol-Sola N., Garcia-Casco A., Proenza J.A. et al. Diamond forms during low pressure serpentinisation of oceanic lithosphere // *Geochemical Perspective Letters*. 2020. Vol. 15. P. 19-24. DOI: 10.7185/geochemlet.2029
15. Simakov S.K., Kouchi A., Mel'nik N.N. et al. Nanodiamond finding in the Hyblean shallow mantle xenoliths // *Scientific Reports*. 2015. Vol. 5. № 10765. DOI: 10.1038/srep10765
16. Romanovsky B. V. Modern catalysis: Science or art? *Soros educational journal*. Volume 6. No. 9,2000, p. 43-48.
17. HETEROGENEOUS CATALYSIS IN PRACTICE, by Charles N. Satterfield, [New York, McGraw-Hill Book Company], 1980. 416 pp. [ISBN] 0-07-054875-F.
18. Robert S. Weber. Perspective: Lies, Damn Lies, and Turnover Rates. *Journal of Catalysis*. Volume 404, December 2021, Pages 925-928. <https://doi.org/10.1016/j.jcat.2021.06.024>
19. Kaplunenko, V., & Kosinov, M. (2025). REVOLUTION IN DIAMOND SYNTHESIS: A unified mechanism of diamond crystallogenesis in nature and in laboratory synthesis. The law of diamond crystallogenesis. *Cambridge Open Engage*. doi:10.33774/coe-2025-56869-v5.
20. Wöhler, F. (1835). *Grundriss der Chemie: Unorganische Chemie* [Foundations of Chemistry: Inorganic Chemistry]. Berlin: Duncker und Humblot. p. 4.
21. Jensen, W. B. (2007). "the origin of the oxidation-state concept". *J. Chem. Educ.* 84 (9): 1418–1419. Bibcode:2007JChEd..84.1418J. doi:10.1021/ed084p1418.
22. William B. Jensen. The Origin of the Oxidation-State Concept. *J. Chem. Educ.* 2007, 84, 9, 1418. <https://doi.org/10.1021/ed084p1418>
23. Viswanathan, B., Gulam Razul, M. Electronegativity provides the relationship between formal charge, oxidation state, and actual charge. *Found Chem* (2022). <https://doi.org/10.1007/s10698-022-09447-6>
24. VALENCE, CHEMICAL BOND, AND EXTENT OF OXIDATION – KEY NOTIONS IN CHEMISTRY Ya. A. UGAI. Угай Я. А. Валентность, химическая связь и степень окисления — важнейшие понятия химии. *Соросовский образовательный журнал*. — 1997. — No 3. — С. 53-57).
25. Nicholas C. 77. Norman, Paul G. Pringle. In defence of oxidation states. *Dalton Transactions* 2022, 51 (2), 400-410. <https://doi.org/10.1039/D0DT03914D>
26. Hans-Peter Looock. Expanded Definition of the Oxidation State. *J. Chem. Educ.* 2011, 88, 3, 282–283. <https://doi.org/10.1021/ed1005213>
27. Swinehart, D.F. More on oxidation numbers. *J. Chem. Educ.* 1952, 29, 284. <https://doi.org/10.1021/ed029p284>
28. R. Resta. Charge States in Transition, *Nature*, 2008, 453, 735.

29. P. Karen , P. McArdle and J. Takats , Towards a Comprehensive Definition of Oxidation State (IUPAC Technical Report), Pure Appl.Chem., 2014, 86 , 1017 —1081. doi:10.1515/pac-2013-0505

30. P. Karen, P. McArdle and J. Takats, Comprehensive Definition of Oxidation State (IUPAC Recommendations 2016), Pure Appl. Chem., 2016, 88, 831 —839.[104] Karen P. Oxidation state, a long-standing issue! Angew Chem Int Ed Engl. 2015 Apr 13;54(16):4716-26. doi: 10.1002/anie.201407561

31. Rebeca G. Castillo, Dr. Anselm W. Hahn, Dr. Benjamin E. Van Kuiken, Dr. Justin T. Henthorn, Jeremy McGale, Prof. Dr. Serena DeBeer. Probing Physical Oxidation State by Resonant X-ray Emission Spectroscopy: Applications to Iron Model Complexes and Nitrogenase. Angewandte Chemie. Volume 60, Issue 18. April 26, 2021. Pages 10112-10121. <https://doi.org/10.1002/anie.202015669>

32. The modern theory of valency. L. Pauling, J. Chem. Soc., 1948, 1461. DOI: 10.1039/JR9480001461.

33. Ning Qin, Sicen Yu, Zongwei Ji, Yanfang Wang, Yingzhi Li, Shuai Gu, Qingmeng Gan, Zhenyu Wang, Zhiqiang Li, Guangfu Luo, Kaili Zhang and Zhongguang Lu. Oxidation State as a Descriptor in Oxygen Reduction Electrocatalysis. CCS Chem. 2022, Just Published. DOI: 10.31635/ccschem.022.202101531. <https://doi.org/10.31635/ccschem.022.202101531>