Chemical-Vapor-Deposited Diamond Film

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February 1999
Chapter 9
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9.1 Introduction

Chemical-vapor-deposited (CVD) diamond film technology looks promising in the field of tribology [9.1–9.5]. CVD diamond coatings offer a broader tribological potential than do natural and high-pressure synthetic diamonds because size and eventually cost will be less of a limitation. In general, however, the advantages and utility of CVD diamond as an industrial ceramic can only be realized if the price is right [9.6]. Until recently skeptics, even internationally well-known tribologists, viewed this technology merely as a rich mother lode of research papers for the academic community. According to Windischmann [9.6] that view may no longer be valid because of two advantages made by a leading CVD diamond supplier in the past year:

1. Reduction of the cost of CVD diamond deposition below $5/carat ($8/cm²)
2. Installation of production capacity

Thus, CVD diamond applications and business in the field of tribology, particularly cutting and forming tools, are an industrial reality [9.1]. At present CVD diamond is produced in the form of coatings or wafers. In this respect it is similar to pyrolytic graphite. Most CVD diamond is produced as films of polycrystalline diamond on ceramic or metal substrates. Coatings with a thickness of submicrometer to 1 mm or more are routinely produced. Thin CVD diamond films are used in film form; thick films are used in a free-standing shape or sheet (wafer) form after removing the substrate. The surface roughness of these coatings is typically in the range 10 nm to 1 μm $R_{a}$ (centerline average) depending on the deposition conditions.

This chapter describes clean and contaminated diamond surfaces, CVD diamond film deposition technology, the characterization of CVD diamond film, and general and tribological properties of these films.
9.2 Diamond Surfaces

9.2.1 Clean Surface

Because diamond has tetrahedral, covalent bonds between each atom and its four nearest neighbors, the free surface may expose dangling bonds. The free surface has high surface energy \( \gamma \), which is associated with dangling bond formation.

Crystalline diamond and silicon all have the diamond structure, where each atom in the crystal is \( sp^3 \) hybridized and tetrahedrally bonded to its four nearest neighbors. When the crystal is cut to expose a crystal plane, chemical bonds are broken along the plane, resulting in half-empty orbitals (dangling bonds) on the surface atoms [9.7, 9.8]. Because of the similarities between the bulk structures of diamond and silicon, a hypothetical surface structure of the silicon crystal, which is the ideal or bulk-terminated structure, is presented in Fig. 9.1 [9.7]. Simple counting of the

![Diagram of silicon surface structures](image)

Figure 9.1.—Dangling bond formation on unreconstructed surfaces of (111), (\( \overline{1}10 \)), and (00\( \overline{1} \)) planes of silicon. (Each silicon atom in the bulk is bonded to its four nearest neighbors in a tetrahedral configuration. The sizes of the silicon atoms are shown to decrease away from the page. From [9.7].)
broken bonds suggests that a surface atom will have two dangling bonds on the 
{001} plane but only one each on the {111} and {100} planes, which are the 
ergetically favored ones. Because of the high energy cost associated with 
dangling bond formation, most diamond surfaces will become reconstructed to 
reduce the number of dangling bonds, thus minimizing their surface free energy. 
The surface valency bonds will back-bond or rehybridize.

Researchers have found that atomically clean diamond has high adhesion and 
friction [9.9, 9.10]. When a fresh atomically clean diamond surface comes in 
contact with an atomically clean surface of counterfacing material, the dangling 
bonds can form strong linkages with bonds on the counterfacing material surface. 
For example, if the surfaces of natural diamond and a metal are cleaned by argon ion 
bombardment, their coefficient of friction is higher than 0.4, as described in 
Chapter 3 and [9.10]. With argon-sputter-cleaned diamond surfaces there are 
probably dangling bonds ready to link up directly with those of another surface.

Because diamond has the highest atomic density of any material, with a molar 
density of 0.293 g-atom/cm³, diamond is the stiffest, hardest, and least compressible of all substances. Tribologically, the real area of contact A is small. Nevertheless, the adhesion and friction of diamond are high in ultrahigh vacuum if the interface is atomically clean and dangling bonds are ready to link up with the counterfacing material surface [9.9, 9.10]. As discussed in Chapter 4, the adhesion and friction of materials are directly related to $sA$ or $\gamma A$. The dangling bonds, high shear strength $s$, and high surface energy $\gamma$ of a clean diamond surface not only provide high friction but also produce high diamond wear due to high adhesion in ultrahigh vacuum. This subject is discussed in Section 9.6.

9.2.2 Contaminated Surface

Although diamond is generally inert to most chemical environments, the interaction of hydrogen and oxygen with diamond surfaces may play an important role in the application of diamond film technology to tribology [9.9-9.11]. A clean diamond surface exposed to atomic hydrogen can adsorb the atomic hydrogen [9.12] and form a hydrogenated diamond surface. Figures 9.2 and 9.3 present a surface structural model for an as-grown {111} diamond surface and an as-grown {100} diamond surface, respectively [9.13]. Almost all carbon atoms on the {111} surface are covered with methyl radical (CH₃) groups. The {100} surface is terminated by the monohydride (CH) group. The chemisorption of hydrogen on the dangling bonds preserves the sp³ hybridization of the surface carbon atoms and stabilizes the diamond structure against the formation of graphite. When a CVD diamond film is annealed to above 1050 °C, the surface hydrogen can be desorbed, destabilizing the diamond surface structure. Consequently, a nondiamond surface layer will be formed on the diamond surface [9.13-9.15].

Significant amounts of oxygen are present on both natural surfaces and those polished in an aqueous medium [9.16]. Oxygen can be chemically bonded to
Figure 9.2.—Surface structural model for diamond \{111\} surface. (The sizes of the carbon and hydrogen atoms are shown to decrease away from the page.) (a) Diamond \{111\} 1x1 with CH$_3$. (b) Diamond \{111\} 1x1 with CH.

diamond surfaces. A hydrogenated diamond surface oxidizes above 300 °C. The oxygen-containing surface species are desorbed as carbon monoxide and carbon dioxide above 480 °C. Both oxygenation and desorption of oxygen-containing surface species simultaneously occur above 480 °C.

Both oxygen-terminated (oxidized) and hydrogen-terminated (hydrogenated) diamond surfaces provide low friction (see Section 9.6 for details). In the atmosphere diamond is known to be one of the slipperiest materials and is similar to polytetrafluoroethylene. This low friction is a surface property that apparently depends on the presence of adsorbed impurities, such as hydrocarbons from the atmosphere, on the oxidized and hydrogenated diamond surfaces.

Both oxidized and hydrogenated diamond surfaces can be fluorinated at temperatures from 300 to 1000 °C when they are treated under carbon tetrafluoride (CF$_4$) plasma conditions [9.16, 9.17]. The C–F bond is observed on CF$_4$-plasma-treated surfaces. Hydrogenated diamond surfaces fluorinate at lower temperatures than oxidized diamond surfaces. However, no fluorination occurs during thermal treatment...
in a CF$_4$ environment. Plasma conditions, using CF$_4$ as a reactant precursor, are necessary for fluorination.

On the other hand, the hydrogenated diamond surface is chlorinated by thermal reaction in chlorine [9.18]. Hydrogen chemisorbed on the diamond surface is abstracted by the chlorine, resulting in chlorine chemisorption. Water vapor can react with the chlorinated diamond surface even at room temperature, producing O–H groups on the surface. A high-temperature treatment of chlorinated diamond with water further oxidizes the diamond surface, producing C=O species on the surface. Water alone, however, cannot oxidize a hydrogenated diamond surface below 500 °C.
Figure 9.4.—Mass deposition rate increase and deposition cost reduction in past decade for CVD diamond produced by direct-current arc jet.
9.3 CVD Diamond Film Deposition Technology

The basic reaction in the chemical vapor deposition of diamond is simple [9.12]. It involves the decomposition of a hydrocarbon, such as methane, as follows:

\[ \text{CH}_4 \rightarrow \text{(Activation)} \rightarrow \text{C(diamond)} + 2\text{H}_2 \]

The carbon species must be activated, since at low pressure graphite is thermodynamically stable and, without activation, only graphite would be formed. Activation is obtained by two basic methods: high temperature and plasma, both of which require a great deal of energy.

Several CVD processes based on these two methods are presently in use. The four most important activation methods at this time are:

1. High-frequency, plasma glow discharge using the microwave and radio-frequency processes
2. Plasma arcing using the direct-current arc and radiofrequency arc processes
3. Thermal CVD using the hot-filament process
4. Combustion synthesis using an oxyacetylene torch

Plasma arcing and combustion synthesis have high substrate deposition rates [9.6, 9.12]. For example, the mass deposition rate using a direct-current arc jet was 20 carats/hr in 1997 (Fig. 9.4(a)). Progress in direct-current arc jet deposition has advanced to the point that the $5/carat (8/cm^2) barrier has been breached (Fig. 9.4(b)). In 1997 diamond made by direct-current arc jet was available at $8/cm^2 (50/in.2).

Diamond has been deposited from a large variety of precursors, including methane, aliphatic and aromatic hydrocarbons, alcohols, ketones, and solid polymers such as polyethylene, polypropylene, and polystyrene [9.12]. These substances generally decompose into two stable primary species: the methyl radicals (CH₃) and acetylene (C₂H₂). The radical is the key compound in generating the growth of CVD diamond.

9.4 Characterization of CVD Diamond

A variety of techniques can be used to characterize CVD diamond films: scanning and transmission electron microscopy (SEM and TEM), to determine surface morphology, microstructure, and grain size; surface profilometry and scanning probe microscopy such as atomic force microscopy (AFM), to measure surface roughness and to determine surface morphology; Rutherford backscattering (RBS) and hydrogen forward scattering (proton recoil detection), to determine the composition including hydrogen; Raman spectroscopy, to characterize atomic bonding state and diamond quality; and x-ray diffraction, to determine the crystal orientation of diamond growth.

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It is generally accepted that, for a material to be recognized as diamond, it must have all of the following characteristics:

1. A clear, sharp diamond peak at 1332 cm\(^{-1}\) in the Raman spectrum
2. A crystalline morphology visible by SEM or TEM
3. A single-phase diamond crystalline structure detectable by x-ray or electron diffraction

Examples as case studies have been prepared focusing attention primarily on microwave-plasma-assisted CVD diamond films and are described in the following subsections.

9.4.1 Surface Morphology and Roughness

The surface morphology and roughness of CVD diamond films can be controlled by varying the deposition parameters, such as gas-phase chemistry parameters and temperatures (e.g., see Table 9.1 and [9.19]). Figure 9.5 shows SEM photomicrographs of fine-, medium-, and coarse-grain diamond films. Triangular crystalline facets typical of diamond are clearly evident on the surfaces of the medium- and coarse-grain diamond films, which have grain sizes estimated at 1100 and 3300 nm, respectively. The grain sizes of the fine-grain diamond were determined from bright- and dark-field electron photomicrographs to be between 20 and 100 nm. The average surface roughness of the diamond films measured by a surface profilometer increases as the grain size increases, as shown in Fig. 9.6.

Note that Fig. 2.20(b) in Chapter 2 is an atomic force microscopy image of a fine-grain CVD diamond film. The fine-grain diamond surface has a granulated or spherulitic morphology (i.e., the surface contains spherical asperities of different sizes).

<table>
<thead>
<tr>
<th>Condition</th>
<th>Substrate*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Si [100]</td>
<td>Si [100]</td>
</tr>
<tr>
<td>Deposition temperature, °C</td>
<td>101±50</td>
</tr>
<tr>
<td>Gaseous flow rate, cm³/min</td>
<td></td>
</tr>
<tr>
<td>CH₄</td>
<td>3.5</td>
</tr>
<tr>
<td>H₂</td>
<td>500</td>
</tr>
<tr>
<td>O₂</td>
<td>0</td>
</tr>
<tr>
<td>Pressure, torr</td>
<td>40</td>
</tr>
<tr>
<td>Microwave power, W</td>
<td>1000</td>
</tr>
<tr>
<td>Deposition time, hr</td>
<td>14</td>
</tr>
<tr>
<td>Thickness, nm</td>
<td>4200</td>
</tr>
<tr>
<td>Grain size, nm</td>
<td>1100</td>
</tr>
<tr>
<td>Surface roughness rms, nm</td>
<td>63</td>
</tr>
</tbody>
</table>

*Scratched with 0.5-μm diamond paste.

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9.4.2 Composition

Figures 9.7 and 9.8 [9,20] present a Rutherford backscattering spectrum and a hydrogen forward-scattering (HFS or proton recoil detection) spectrum, respectively, of the fine-grain CVD diamond film shown in Fig. 9.5(a). Besides carbon from the diamond film and silicon from the substrate, no other elements are
observed in the RBS spectrum. From both the RBS and HFS spectra it is estimated that the fine-grain diamond film consists of 97.5 at.% carbon and 2.5 at.% hydrogen. (In contrast, the medium- and coarse-grain diamond films contained less than 1 at.% hydrogen [9.19].) Both carbon and hydrogen are uniformly distributed in the fine-grain diamond film from the top of the surface to the silicon substrate.

The RBS analytical results can also be used to determine diamond film thickness. Figure 9.7 also presents a simulated RBS spectrum of the fine-grain diamond film containing a carbon-to-hydrogen ratio (C/H) of 97.5/2.5 obtained by using the RUMP computer code [9.21]. In the computer program the thickness of the diamond film is taken as a variable and is obtained from the close match between the observed and simulated RBS, as shown in Fig. 9.7. The film thickness of the fine-grain CVD diamond is 1.5 μm at the center of the substrate, and the deposition rate is estimated to be 0.14 μm/hr.
Figure 9.7.—Rutherford backscattering spectrum of fine-grain diamond film on silicon substrate. (Simulation curve was calculated by using the computer code RUMP.)

Figure 9.8.—Hydrogen forward-scattering spectrum (proton recoil analysis) of fine-grain diamond film on silicon substrate.
9.4.3 Atomic Bonding State

Figure 9.9 shows typical Raman spectra of the fine- and medium-grain diamond films. Both spectra show one Raman band centered at 1332 cm\(^{-1}\) and one at 1500 to 1530 cm\(^{-1}\). The sharp peak at 1332 cm\(^{-1}\) is characteristic of the sp\(^3\) bonding of the diamond form of carbon in the film. The very broad peak centered around 1530 cm\(^{-1}\) is attributed to the sp\(^2\) bonding of the nondiamond forms of carbon (graphite and other forms of carbon) [9.22–9.24].

More diamond was produced in the medium-grain CVD diamond films (e.g., Fig. 9.9) than in the fine-grain films, as is evident from the relative intensities of the diamond and nondiamond carbon Raman bands [9.19]. However, the ratio of the intensities of the Raman responses at 1332 and 1530 cm\(^{-1}\) does not indicate the ratio of diamond to nondiamond carbon present in the film. The Raman technique is approximately 50 times more sensitive to sp\(^2\)-bonded (nondiamond) carbon than it is to sp\(^3\)-bonded (diamond) carbon [9.23]. Thus, the peak at approximately 1530 cm\(^{-1}\) for each film represents a much smaller amount of nondiamond carbon in these diamond films. The Raman spectrum of the coarse-grain diamond film was similar to that of the medium-grain diamond film.

9.4.4 Microstructure

X-ray diffraction analysis (XRD) is used to determine the structure and crystal orientation of the CVD diamond films. Figure 9.10 shows typical XRD patterns for the fine- and medium-grain diamond films [9.19]. Peaks representing only the diamond film and the silicon substrate appear in the XRD spectra. Diffraction peaks corresponding to the \{111\}, \{220\}, \{311\}, and \{400\} planes, reflective of diamond, are clearly evident. The intensity ratios I\{220\}/I\{111\} were calculated from the XRD patterns for the fine- and medium-grain diamond films and found to be 1.3 and 0.04, respectively. The powder diffraction pattern of diamond with random crystal orientation (ASTM 6-0675) gives I\{220\}/I\{111\} = 0.27.

Thus, most crystallites in the fine-grain diamond film are oriented along the (110) direction. Other researchers [9.22] have observed similar [110] crystal orientation texture in microcrystalline diamond film grown in a \{100\} silicon substrate by an activated CVD technique. Most crystallites in the medium-grain diamond films are oriented along the \{111\} direction, and the \{111\} planes are parallel to the surface. The well-formed triangular facets observed in SEM photomicrographs (Fig. 9.5) of the medium- and coarse-grain diamond films confirm the \{111\} crystal orientation.

Figure 9.11 presents the transmission electron diffraction pattern, a transmission electron bright-field photomicrograph, and a transmission electron dark-field photomicrograph of the free-standing, fine-grain CVD diamond film [9.20]. Diffraction rings and dots are observed in the selected-area diffraction (SAD) pattern (Fig. 9.11(a)). The d spacings of the diffraction rings were calculated by using an
Figure 9.9.—Raman spectra of diamond films. (a) Fine-grain (20 to 100 nm) diamond film on {100} silicon substrate. (b) Medium-grain (1100 nm) diamond film on {100} silicon substrate.
Figure 9.10.—X-ray diffraction patterns of diamond films. (a) Fine-grain (20 to 100 nm) diamond film on {100} silicon substrate. (b) Medium-grain (1100 nm) diamond film on {100} silicon substrate.
Figure 9.11.—Free-standing diamond films.
(a) Selected-area diffraction pattern.
(b) Bright-field TEM. (c) Dark-field TEM.
aluminum SAD pattern as a calibration standard and found to match well with the known diamond d spacings. No evidence of nondiamond carbon is found in the SAD pattern. This observation indicates that the concentration of nondiamond component in the diamond film is very small.

Careful observation of the bright-field photomicrograph (Fig. 9.11(b)) shows various nuclei-like regions marked N. Diamond grains are distributed radially outward from these nuclei. A grain boundary is formed where the grains from various nuclei meet. The grain sizes were estimated from the dark-field photomicrograph (Fig. 9.11(c)) and found to vary from 20 to 100 nm.

### 9.5 General Properties of CVD Diamond

Diamond consists of light carbon atoms held together by strong forces, and this combination produces many extreme properties. Table 9.2 summarizes the general properties of CVD diamond and compares them to those of single-crystal diamond, such as type Ib. This table and the description in this section are based on a table from Pierson [9.12] and other references [9.25–9.27]. Because of the difficulty in testing, the effect of impurities and structural defects, and the differences between the various deposition processes, uncertainty and spread have been found in the reported property values. The values in Table 9.2 should be viewed with caution.
### TABLE 9.2.—PROPERTIES OF CVD DIAMOND

<table>
<thead>
<tr>
<th>Property</th>
<th>CVD diamond</th>
<th>Single-crystal diamond</th>
</tr>
</thead>
<tbody>
<tr>
<td>Density, g/cm³</td>
<td>3.51</td>
<td>3.515</td>
</tr>
<tr>
<td>Thermal conductivity at 25 °C, W/m²°C⁻¹</td>
<td>2100</td>
<td>2200</td>
</tr>
<tr>
<td>Thermal expansion coefficient at 25 to 200 °C, °C⁻¹</td>
<td>2.0×10⁹</td>
<td>(1.5 to 4.8)×10⁹</td>
</tr>
<tr>
<td>Tensile strength, GPa</td>
<td>1 to 5</td>
<td>4 to 6</td>
</tr>
<tr>
<td>Young's modulus, Pa</td>
<td>(7 to 9)×10¹¹</td>
<td>10.5×10¹¹</td>
</tr>
<tr>
<td>Poisson’s ratio, GPa</td>
<td>0.1 or 0.07</td>
<td>0.1 or 0.07</td>
</tr>
<tr>
<td>Vicker’s hardness range, kg/nm²</td>
<td>5000 to 10000</td>
<td>5700 to 10400</td>
</tr>
</tbody>
</table>

**Chemical properties**

Both CVD and single-crystal diamonds are resistant to all liquid organic and inorganic acids, alkalis, and solvents at room temperature. One of the most chemically resistant materials.

<table>
<thead>
<tr>
<th>Property</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Band gap, eV</td>
<td>5.45</td>
<td>5.45</td>
</tr>
<tr>
<td>Index of refraction at 10 μm</td>
<td>2.34 to 2.42</td>
<td>2.40</td>
</tr>
<tr>
<td>Electrical resistivity, ohm-cm</td>
<td>10¹⁰ to 10¹⁰</td>
<td>10¹⁰</td>
</tr>
<tr>
<td>Dielectric constant (45MHz to 20 GHz)</td>
<td>56</td>
<td>5.70</td>
</tr>
<tr>
<td>Dielectric strength, V/cm</td>
<td>10¹⁰</td>
<td>10¹⁰</td>
</tr>
<tr>
<td>Loss tangent (45 MHz to 20 GHz)</td>
<td>&lt; 0.0001</td>
<td></td>
</tr>
<tr>
<td>Saturated electron velocity</td>
<td>27</td>
<td></td>
</tr>
<tr>
<td>Carrier mobility:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Electron (n)</td>
<td>1350 to 1500</td>
<td>2200</td>
</tr>
<tr>
<td>Positive hole (p)</td>
<td>480</td>
<td>1600</td>
</tr>
</tbody>
</table>

*Varies with crystal orientation (see Chapter 8).

In diamond the light carbon atoms are densely packed, but the lattice is in many ways similar to those of other crystals, such as silicon, of the same structure. Therefore, diamond is a lightweight material, the density of both CVD diamond and single-crystal diamond being approximately 3.5 g/cm³ (Table 9.2). Note that the closeness of carbon atoms leads to a small compressibility, as described in Chapter 8, and the specific heat is also small (6.19 J g-atom⁻¹°C⁻¹) at room temperature.

One of the many remarkable properties of diamond is its high thermal conductivity, as shown in Fig. 9.12. The data in this figure were taken from Pierson [9.12] and Windischmann [9.6]. For the highest quality single-crystal, type IIa diamonds, the thermal conductivity is about 25 W cm⁻¹°C⁻¹ at room temperature, or more than six times that of copper (4 W cm⁻¹°C⁻¹). Such high thermal conductivity is caused by the stiffness of the diamond bond and the diamond structure. Other types of single-crystal diamond and CVD diamond have lower thermal conductivity (Table 9.2) at room temperature because they contain larger concentrations of impurities than does type IIa diamond.

Using values of 10.5×10¹¹ Pa, 5.5×10¹¹ Pa, 5.3 J/m, and 0.154 nm for the Young’s modulus, the shear modulus, the fracture surface energy, and the nearest neighbor distance, respectively, the theoretical strength in tension of single-crystal diamond is 1.9×10¹¹ Pa in the (111) direction (i.e., E/5, where E is the modulus of elasticity [9.25]). Theoretical strength in shear equals 12.1×10¹⁰ Pa in the (110) [111] system.
(i.e., \(-G/4\), where \(G\) is the shear modulus). Actual tensile strengths for the octahedral \([111]\) and cubic \([100]\) planes of single-crystal type IIa diamond are 3.8 and 3.7 GPa, respectively. The average is \(\sim 3.75\) GPa for a Poisson’s ratio of 0.1 (or \(\sim 4.0\) GPa for a Poisson’s ratio of 0.07), roughly 1/50th of the theoretical strength. Actual tensile strengths obtained by many workers range from 4 to 6 GPa for single-crystal diamond and from 1 to 5 GPa for CVD diamond (Table 9.2). Many workers have found Young’s moduli for CVD diamond in the range \(7 \times 10^{11}\) to \(9 \times 10^{11}\) Pa and Poisson’s ratios of 0.1 or 0.07 [9.25–9.27].

Chapter 8 states that diamond is the hardest known material and that the hardness of single-crystal diamond varies by a factor 2 or more as a function of crystal orientation and impurities. This statement is also true for CVD diamond. Figure 9.13, based on a figure from [9.12] and other references [9.25–9.30], shows the high indentation hardness for diamond as compared with a few other hard materials.

Diamond is chemically resistant to all liquid organic and inorganic acids at room temperature. However, it can be etched by several compounds, including strong oxidizers such as sodium and potassium nitrates above 500 °C, by fluxes of sodium and potassium chlorates, and by molten hydroxides, such as sodium hydroxide (NaOH). Diamond is resistant to alkalis and solvents. At approximately 1000 °C it reacts readily with carbide-forming metals such as iron, cobalt, nickel, aluminum, tantalum, and boron. Generally speaking, diamond can be considered as one of the most chemical-resistant materials. The chemical properties of CVD diamond are similar to those of single-crystal diamond.
Figure 9.13.—Indentation hardness of diamond and other hard materials.

Table 9.2 also includes optical, electrical, and semiconductor properties of diamond as a reference.

9.6 Friction and Wear Properties of CVD Diamond

9.6.1 Humid Air and Dry Nitrogen Environments

The tribological properties of CVD diamond are similar to those of natural and synthetic diamond. The coefficient of friction and wear resistance of CVD diamond are generally superior in the atmosphere. However, the environment to which a CVD diamond film is exposed can markedly affect its tribological properties, such as friction and wear behavior. They vary with the environment, possessing a Jekyll-and-Hyde character [9.19].

When the fine-, medium-, and coarse-grain diamond films characterized in Section 9.4 were brought into contact with a natural diamond pin in reciprocating sliding motion in air and in nitrogen, the coefficients of friction varied as the pin traveled back and forth (reciprocating motion), retracing its tracks on the diamond films (Fig. 9.14).
Figure 9.14.—Coefficient of friction as function of number of passes of bulk diamond pin in contact with fine-, medium-, and coarse-grain diamond films (a) in humid air (approx. 40% relative humidity) and (b) in dry nitrogen.
Both in humid air at a relative humidity of 40% and in dry nitrogen, abrasion occurred and dominated the friction and wear behavior. The bulk natural diamond pin tended to dig into the surface of diamond films during sliding and produce a wear track (groove). SEM observations of the diamond films indicated that small fragments chipped off their surfaces. When abrasive interactions between the diamond pin surface and the initially sharp tips of asperities on the diamond film surfaces were strong, the friction was high (points A in Fig. 9.14). The surface roughness of diamond films can have an appreciable influence on their initial friction (i.e., the greater the initial surface roughness, the higher the initial coefficient of friction, Fig. 9.15(a)). Similar frictional results have also been found by other workers on single-crystal diamonds [9.31] and on diamond coatings [9.32–9.34].

As sliding continued and the pin passed repeatedly over the same track, the coefficient of friction was appreciably affected by the wear on the diamond films (Fig. 9.14) (i.e., a blunting of the tips of asperities). When repeated sliding produced a smooth groove or a groove with blunted asperities on the diamond surface (Fig. 9.16), the coefficient of friction was low, and the initial surface roughness effect became negligible. Therefore, the equilibrium coefficient of friction was independent of the initial surface roughness of the diamond film (Fig. 9.15(b)).

The generally accepted wear mechanism for diamonds is that of small fragments chipping off the surface [9.9, 9.19]. This mechanism is in agreement with the wear of diamond films. The wear rate is dependent on the initial surface roughness of the diamond films (Fig. 9.17), increasing markedly with an increase in initial surface roughness. The wear rates of the diamond films in humid air and in dry nitrogen are comparable to the wear rates of single-crystal diamonds and diamond films investigated by other workers [9.32, 9.35, 9.36].

9.6.2 Ultra-High-Vacuum Environment

When the fine-, medium-, and coarse-grain diamond films were brought into contact with a natural diamond pin in unidirectional pin-on-disk sliding motion in vacuum, the coefficients of friction were high and varied with the number of passes [9.15]. In vacuum, as in humid air and in dry nitrogen, the bulk natural diamond pin dug into the surfaces of the diamond films during sliding and produced a wear track (groove, Fig. 9.18). The groove surface was generally smoother than the original surface of the diamond films. Further analysis of the grooves by scanning electron microscopy revealed that the tips of the diamond coating asperities were worn smooth and that the gaps between asperities were filled by debris.

The coefficient of friction increased with an increase in the number of passes (Fig. 9.19), just the opposite of what occurred in humid air and in dry nitrogen. Further, the initial surface roughness of the diamond film had no effect on friction. These results lead us to ask the following questions: What factors determine friction behavior? Have dangling bonds been exposed during sliding and played a role in the
Figure 9.15.—Coefficient of friction as function of initial surface roughness of diamond films in humid air (approx. 40% relative humidity) and in dry nitrogen. (a) Initial coefficients of friction. (b) Equilibrium coefficients of friction.
Figure 9.16.—Wear tracks (grooves) on diamond films after 30,000 passes of bulk diamond pin in dry nitrogen. (a) Fine-grain (20 to 100 nm) diamond film; rms surface roughness, 15 nm. (b) Medium-grain (1100 nm) diamond film; rms surface roughness, 63 nm. (c) Coarse-grain (3300 nm) diamond film; rms surface roughness, 160 nm.
Figure 9.17.—Wear rates of diamond films as function of diamond surface roughness in humid air and in dry nitrogen.
Figure 9.18.—Wear tracks (grooves) on diamond films after 100 passes of bulk diamond pin in ultrahigh vacuum. (a) Fine-grain (20 to 100 nm) diamond film on silicon substrate; rms surface roughness, 15 nm. (b) Medium-grain (1000 nm) diamond film on silicon nitride substrate; rms surface roughness, 52 nm. (c) Coarse-grain (1500 nm) diamond film on α-silicon carbide substrate; rms surface roughness, 92 nm.

Figure 9.19.—Typical friction trace for bulk diamond pin in contact with diamond film on α-silicon carbide substrate in ultrahigh vacuum (initial coefficient of friction, $\mu_I$; equilibrium coefficient of friction, $\mu_F$).
friction behavior? Which is more important for diamond surfaces in vacuum, abrasion or adhesion?

Removing some of the contaminant surface film from the contact area of diamond films by sliding action resulted in stronger interfacial adhesion between the diamond pin and the diamond films and raised the coefficient of friction, as shown in Fig. 9.19. A contaminant surface film may be removed by repeatedly sliding the diamond pin over the same track in ultrahigh vacuum [9.36].

The friction results shown in Fig. 9.19 are in agreement with other researchers' results for single-crystal diamond rubbing against diamond and for CVD diamond sliding against CVD diamond in vacuum [9.37, 9.38]. At a pressure of 93 nPa Bowden and Hanwell [9.38] observed an initial coefficient of friction of 0.1 for diamond on diamond; within several hundred passes, however, the coefficient of friction rose rapidly to 0.9 and remained constant. Dugger, Peebles, and Pope [9.39] also found that the coefficient of friction increased to 0.47 when CVD diamond slid against itself in vacuum (<0.6 μPa). In both cases the increase in friction was attributed to cleaning the adsorbed contaminants from the surface by rubbing or sliding in vacuum at room temperature.

When sliding continues, the wear dulls the tips of the diamond grains and increases the contact area in the wear track, thereby causing an increase in friction. The increase in equilibrium friction that results from cleaning off the contaminant surface film by sliding and from increasing the contact area is greater than the corresponding decrease in abrasion and friction that results from blunting the tips of surface asperities. This relationship is brought out clearly in Fig. 9.20; here the equilibrium coefficients of friction (1.5 to 1.8) are greater than the initial coefficients of friction (1.1 to 1.3) regardless of the initial surface roughness of the diamond films. In vacuum, therefore, the friction arises primarily from adhesion between the sliding surfaces of the diamond pin and the diamond films.

The wear rates of the diamond films in ultrahigh vacuum (Fig. 9.21) depended on the initial surface roughness of the diamond films, generally increasing with an increase in initial surface roughness. The wear rates of the diamond films in ultrahigh vacuum were considerably higher than those of the diamond films in humid air or in dry nitrogen (Fig. 9.17). Obviously, under these vacuum conditions adhesion between the sliding surfaces of the diamond pin and the diamond film plays an important role in the higher wear process.

Thus, under vacuum conditions it is adhesion between the sliding surfaces of the diamond pin and diamond films (due to the highly clean state) and the possible presence of dangling bonds that play a significant role in the friction and wear process. The surface roughness of the diamond films does not have much influence on the friction of diamond films in ultrahigh vacuum.
Figure 9.20.—Initial ($\mu_I$) and equilibrium ($\mu_F$) coefficients of friction as function of initial surface roughness of diamond films in ultrahigh vacuum.

Figure 9.21.—Wear factors of diamond films in ultrahigh vacuum as function of initial surface roughness.
References


### 13. ABSTRACT (Maximum 200 words)

This chapter describes the nature of clean and contaminated diamond surfaces, CVD diamond film deposition technology, analytical techniques and the results of research on CVD diamond films, and the general properties of CVD diamond films. Further, it describes the friction and wear properties of CVD diamond films in the atmosphere, in a controlled nitrogen environment, and in an ultra-high-vacuum environment.