Effects of Air Exposure on Hard and Soft X-ray Photoemission Spectra of Ultrananocrystalline Diamond/Amorphous Carbon Composite Films

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Abstract: Hard X-ray photoemission spectroscopy (HAXPES) was employed for the structural evaluation of ultrananocrystalline diamond/amorphous carbon (UNCD/a-C) composite films deposited on cemented carbide substrates, at substrate temperatures up to 550 °C by coaxial arc plasma deposition. The results were compared with those of soft X-ray photoemission spectroscopy (SXPES). Since nanocrystalline diamond grains are easily destroyed by argon ion bombardment, the structural evaluation of UNCD/a-C films, without the argon ion bombardment, is preferable for precise evaluation. For samples that were preserved in a vacuum box after film preparation, the sp³ fraction estimated from HAXPES is in good agreement with that of SXPES. The substrate temperature dependencies also exhibited good correspondence with that of hardness and Young’s modulus of the films. On the other hand, the sp³ fraction estimated from SXPES for samples that were not preserved in the vacuum box had an apparent deviation from those of HAXPES. Since it is possible for HAXPES to precisely estimate the sp³ fraction without the ion bombardment treatment, HAXPES is a feasible method for UNCD/a-C films, comprising nanocrystalline diamond grains.

Keywords: atomic structures; HAXPES; hardness; ion bombardment; nanocomposite coating; UNCD/a-C films

1. Introduction

Nanocrystalline diamond (NCD) films have shown promise as a hard coating material for enhancing the performance of cutting tools and prolonging tool lifetime. NCD films introduce the highest known hardness, a smooth surface, and a low friction coefficient, resulting in fine diamond grains, less than 100 nm in size [1–3]. Ultrananocrystalline diamond (UNCD) films, a kind of NCD film comprising grains less than 10 nm [4,5], are a promising candidate as a hard coating application on cutting tools [6].
Precisely, UNCD films comprise a large number of nano-sized diamond grains, an amorphous carbon matrix [7], and a large number of structurally specific grain boundaries in the films. They are prepared by microwave plasma CVD [8,9] and coaxial arc plasma deposition (CAPD) [10–14]. When CAPD is used, film deposition is possible in hydrogen-free atmospheres, which results in films containing a greater amount of nonhydrogenated amorphous carbon (a-C), compared with those prepared by CVD. Thus, when prepared with CAPD, we distinctively describe them as UNCD/a-C films.

It is well known that cathodic arc discharge deposition makes possible the growth of hydrogen-free sp³-rich a-C films [15]. In addition, CAPD also has the unique feature where a coaxial arc plasma gun equipped with an anodic cylinder can bunch ions ejected from a cathodic graphite rod located in the cylinder. Owing to this structure, a supersaturated condition with highly energetic carbon ions, desired for the growth of UNCD crystallites [14], can be realized on substrates [16,17]. The high hardness of UNCD/a-C films is attributed to the existence of UNCD crystallites and the sp³-rich a-C matrix [18].

Careful characterization of the atomic structures is essential since the sp³ fraction determines the mechanical properties of carbon films. The sp³ fraction is estimated by various methods, such as Raman spectroscopy, soft and hard X-ray photoemission spectroscopy, Auger electron spectroscopy, near edge X-ray absorption fine structure (NEXAFS), and high energy electron loss spectroscopy (HEELS).

Among the methods of sp³ characterization, Raman spectroscopy is a popular optical technique used to investigate bonding structures of carbon films. Although, there has been a debate regarding the accuracy of identifying the UNCD crystallites using this method [19,20] and it can’t estimate the sp³ fraction precisely [21]. While the NEXAFS and HEELS techniques are available for analyzing bonding structures, they also show some uncertainty in estimating the sp³ fraction [22].

For the X-ray photoemission spectroscopic characterization of UNCD/a-C films, the influence of surface condition of the films spectra, is not negligible [23]. The soft X-ray photoemission spectroscopy (SXPES), employed mainly for investigating the atomic structures of carbon films [24], detects photoelectrons ejected from surfaces a few nanometers in depth. Thus, to exclude the influence of surface conditions, surface contaminants are removed with ion bombardments before measurement. When applied to UNCD/a-C films, nano-sized diamond crystallites are easily destroyed by the bombardment of Argon ions and it is therefore difficult to evaluate the original structure around the film surfaces [23].

Hard X-ray photoemission spectroscopy (HAXPES) has attracted attention in recent years as an alternative technique that solves the above-mentioned problem, as it examines buried regions at depths of more than 10 nm by employing hard X-ray incidence beams. In other words, it detects spectra mainly from bulk electronic structures on the films subsurface [25,26]. In this work, HAXPES was applied to the structural evaluation of UNCD/a-C films and correlations between the mechanical and structural properties were studied. In addition, HAXPES results were compared to those of SXPES and the feasibility of HAXPES for the evaluation of UNCD/a-C films was investigated.

2. Materials and Methods

2.1. Film Preparation

UNCD/a-C films were deposited on WC-Co substrates (K-type cemented tungsten carbide) with dimensions of φ10 × 4.5 mm by CAPD, employing a coaxial arc plasma gun (APG-1000, ULVAC, kanagawa, Japan) equipped with a graphite rod (purity of 99.99%, φ10 × 30 mm). The pretreatment of the WC-Co substrates by chemical etching for removing Co near the surface, which has been indispensably applied for the hard coating of polycrystalline diamond on WC-Co by CVD, was not carried out in this study. Only the pretreatment of roughening the WC-Co surface was made in the range of Ra = 0.15 µm to 0.2 µm, by dissolving the WC using chemical etching to enhance adhesion of the films. Before the deposition of the films, the inside of the vacuum chamber was evacuated to be
less than 10^{-4} \text{Pa} by a turbomolecular pump. A voltage of 100 \text{V} was applied to the coaxial arc plasma gun equipped with a 720 \mu \text{F} capacitor and the arc discharge of the arc plasma gun was operated at a repetition rate of 1 \text{Hz}. The films were deposited at different substrate temperatures ranging from room temperature to 550 °C. The details of film preparation conditions are presented in our previous papers [15,16].

2.2. Film Characterization

The samples were divided into two groups according to a difference in the storage procedure after deposition. The difference is schematically shown in Figure 1. The samples of Group I were exposed naturally to surrounding air for a maximum of two weeks after finishing deposition by CAPD apparatus and until HAXPES and SXPES measurements were made. On the other hand, the samples of Group II were preserved in a portable vacuum box containing silica gel, where the inside pressure is less than 0.1 \text{MPa}, as soon as they were taken out of the CAPD apparatus, for a maximum of two weeks. Moreover, they were quickly set into an SXPES vacuum apparatus before measurements were taken.

The atomic structures of the films were examined by HAXPES and SXPES. While, HAXPES was applied to the samples of Group I, SXPES investigated the samples of Group I and II. The chemical compositions and sp^3/(sp^3 + sp^2) ratio were investigated from SXPES and HAXPES spectra. The HAXPES measurements with an incident photon energy of 8 \text{keV} were conducted at BL47XU in Spring-8. SXPES measurements with an incident photon energy of 350 eV were performed at BL12 of SAGA Light Source/Kyushu Synchrotron Light Source Center. In addition, SXPES wide-range measurements with a photon energy of 1253.6 eV (Mg Kα line) were made for investigating the chemical composition of the films. The spectra were decomposed into peaks using a Voigt function (80% Gaussian and 20% Lorentzian) after subtracting backgrounds [27].

Each sample was tested ten times to calculate mean values of hardness and Young’s modulus by nano-indentation test at an applied load of 0.5 \text{mN} (HM500, Fisher Instruments Company, Sindelfingen, Germany).

3. Results and Discussions

Table 1 indicates the hardness and Young’s modulus of films deposited at different substrate temperatures. The hardness and Young’s modulus decrease with increasing substrate temperature,
which implies that the atomic structures of films are affected by the increase in substrate temperature. To study the origin, SXPES and HAXPES were employed.

Table 1. Typical preparation conditions of the UNCD/a-C films.

<table>
<thead>
<tr>
<th>Preparation Conditions</th>
<th>Sample 1</th>
<th>Sample 2</th>
<th>Sample 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Substrate temp. (°C)</td>
<td>Room temp. (R.T.)</td>
<td>300</td>
<td>550</td>
</tr>
<tr>
<td>Deposition rate (nm/pulse)</td>
<td>0.29</td>
<td>0.35</td>
<td>0.35</td>
</tr>
<tr>
<td>Hardness (GPa)</td>
<td>51.3</td>
<td>21.2</td>
<td>11.7</td>
</tr>
<tr>
<td>Young’s Modulus (GPa)</td>
<td>520.2</td>
<td>271.1</td>
<td>121.9</td>
</tr>
</tbody>
</table>

Figure 2 shows the SXPES and HAXPES survey-spectra. The spectra do not show any peaks due to substrate elements, which means that the diffusion from substrates into films rarely occurs, even at the highest substrate temperature of 550 °C [15]. They show only significant intense C 1s peaks and weak O 1s peaks from collected photoelectrons of excited core levels of carbon and oxygen elements. The O 1s peak at 532 eV is mainly ascribed to the adsorbed oxygen on the surface of the film [28]. The HAXPES O 1s peaks are obviously weak comparing with those of the SXPES O 1s peaks. In addition, the HAXPES detects spectra mainly from bulk electronic structures at buried layers of the films [29]. This means that the O 1s peak is partially attributed to residual oxygen in the chamber, resulted from water vapor contamination during the films deposition.

Figure 3 shows the C 1s spectra of Group I samples (a), (b), and (c), measured by SXPES and (d), (e), and (f), measured by HAXPES, and Group II samples (g), (h), and (i), measured by SXPES. Macroscopically, the maximum energy of the spectra shifts to a higher binding energy with decreasing substrate temperature, which is particularly evident for Group I spectra measured by HAXPES and Group II spectra measured by SXPES. The shifts might have occurred due to the enhancement of the sp<sup>3</sup> bonding fraction [30], which is consistent with hardness increasing as substrate temperature decreases.

**Figure 2.** Soft X-ray photoemission spectroscopic (SXPES) survey (a–d), and hard X-ray photoemission spectroscopic (HAXPES) survey (e–h), of Group I ultrananocrystalline diamond/amorphous carbon (UNCD/a-C) films deposited at substrate temperature of room temperature (R.T.), 300 °C, and 550 °C, and WC-Co substrate.
The HAXPES measurements show sharp spectra with very weak shoulder of the C–O/C–O–C peak, profile of the sp^3 peaks should be ascribed to the adsorption of oxygen on the film surface during and after films deposition [23,33]. The C=O/COOH peak is observed in the spectra measured by SXPES for Group I samples. The Group II samples preserved in the vacuum box produced relatively consistent peaks, compared to those revealed by HAXPES. Figure 4 shows peaks, unlike those obtained by SXPES for Group I samples. The Group II samples preserved in the vacuum box. The vacuum box contains powdered silica gel that absorbs humidity and keeps the inside of the vacuum box dry. This means that the contribution of contaminants found in the upper first layers of the films is limited, due to preserving the samples in the portable vacuum box. The HAXPES measurements show sharp spectra with very weak shoulder of the C–O/C–O–C peak, due to investigation at deep depth in the films. However, the argon ion bombardment is essential for cleaning the films surface prior to X-ray photoemission spectroscopic measurements, to reduce the contribution of film surface contaminants. The HAXPES method makes possible the structural analysis of Group II samples which means the adsorption of oxygen is suppressed, owing to their preservation in the portable vacuum box. The vacuum box contains powdered silica gel that absorbs humidity and keeps the inside of the vacuum box dry. This means that the contribution of contaminants found in the upper first layers of the films is limited, due to preserving the samples in the vacuum box. The HAXPES measurements show sharp spectra with very weak shoulder of the C–O/C–O–C peak, due to investigation at deep depth in the films. However, the argon ion bombardment is essential for cleaning the films surface prior to X-ray photoemission spectroscopic measurements, to reduce the contribution of film surface contaminants. The HAXPES method makes possible the structural examination of the UNCD/a-C films without applying any surface pretreatment, such as argon ion bombardment, which destroys nano-sized diamond crystallite in the films [23].

The sp^3 peaks show a relative decrease as substrate temperature increases. The HAXPES technique produces sharp and intense sp^3 peaks with a noticeable difference compared to the sp^2 peaks, unlike those obtained by SXPES for Group I samples. The Group II samples preserved in the vacuum box produced relatively consistent peaks, compared to those revealed by HAXPES. Figure 4 shows the substrate temperature dependence of the hardness and sp^3 bonding fraction. The dependence profile of the sp^3 bonding fraction estimated from HAXPES and SXPES for Group II samples are in good agreement, not only with each other but also with the hardness dependence profile. This means that the estimation of the sp^3 fraction is precise, since the hardness proportionally increases with the sp^3 fraction [34,35]. On the other hand, the substrate temperature dependence profile of the sp^3 fraction estimated from SXPES for Group I, obviously deviates from that of hardness. Estimation of the sp^3 bonding fraction is also affected by the existence of C–O/C–O–C and C=O/COOH peaks. The SXPES investigation for Group I emphasizes that the presence of undesirable contaminants on the films surface could lead to difficulties in obtaining surface functionalization. Therefore, the HAXPES...
method and demonstrating a careful methodology for SXPES investigation, are recommended when assessing the sp³ bonding fraction to UNCD/a-C films and nanomaterials.

4. Conclusions

The atomic structures of UNCD/a-C composite films were investigated by HAXPES and SXPES. It was found that the sp³ fraction estimated from HAXPES spectra corresponds well with the hardness and Young’s modulus in their substrate temperature dependencies. On the other hand, the sp³ fraction estimated from SXPES spectra of the samples that were exposed to air after film preparation deviated from that of HAXPES, while SXPES results of samples that were preserved in the vacuum box were in good agreement with those of HAXPES. This indicates that it is possible for HAXPES to precisely estimate the sp³ fraction and similarly to SXPES when samples are preserved in a vacuum box. HAXPES is a promising technique for structural characterization of UNCD/a-C films because the condition of the film surface has a negligible effect on the spectra of films.


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Conflicts of Interest: The authors declare no conflict of interest.

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