Contactless Photoconductance Study on Undoped and Doped Nanocrystalline Diamond Films


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ABSTRACT: Hydrogen and oxygen surface-terminated nanocrystalline diamond (NCD) films are studied by the contactless time-resolved microwave conductivity (TRMC) technique and X-ray photoelectron spectroscopy (XPS). The optoelectronic properties of undoped NCD films are strongly affected by the type of surface termination. Upon changing the surface termination from oxygen to hydrogen, the TRMC signal rises dramatically. For an estimated quantum yield of 1 for sub-bandgap optical excitation the hole mobility of the hydrogen-terminated undoped NCD was found to be ∼0.27 cm²/(V s) with a lifetime exceeding 1 μs. Assuming a similar mobility for the oxygen-terminated undoped NCD a lifetime of ∼100 ps was derived. Analysis of the valence band spectra obtained by XPS suggests that upon oxidation of undoped NCD the surface Fermi level shifts (toward an increased work function). This shift originates from the size and direction of the electronic dipole moment of the surface atoms, and leads to different types of band bending at the diamond/air interface in the presence of a water film. In the case of boron-doped NCD no shift of the work function is observed, which can be rationalized by pinning of the Fermi level. This is confirmed by TRMC results of boron-doped NCD, which show no dependency on the surface termination. We suggest that photoexcited electrons in boron-doped NCD occupy nonionized boron dopants, leaving relatively long-lived mobile holes in the valence band.

KEYWORDS: photoconductance, nanocrystalline diamond, TRMC, carrier lifetime, mobility, contactless

I. INTRODUCTION

Since the past decade, the interest in diamond research, especially in nanocrystalline diamond (NCD) films, has grown significantly because of its unique material properties. NCD films exhibit high chemical stability in harsh environments, allow biochemical surface functionalization, and can be doped with impurities to alter its electronic properties. Furthermore, NCD films can be fabricated on wafer-scale at low-cost using the chemical vapor deposition (CVD) technique. As a result, NCD films could be a solution to the growing need for a better understanding of the factors affecting the optoelectronic properties to further control and improve the film quality. Several studies on the defect distribution in the microcrystalline and NCD films using photoluminescence techniques have been reported. In addition, the influence of experimental conditions such as ambient temperature, ultraviolet light irradiation, and air pressure on photoexcited charge characteristics using photoluminescence techniques have also been discussed. Although photoluminescence techniques give important information on charge carrier dynamics of sub-bandgap states in NCD films, it is found to be less sensitive to the surface termination effect. Additionally, the surface properties of NCD films are significantly affected by hydrogen and oxygen termination in terms of surface wettability, electrical con-
ductivity and electron affinity.\textsuperscript{18,19} Therefore, it is imperative to study the effect of surface termination on the charge carrier kinetics and degree of Fermi level pinning using a complementary technique that can investigate surface sensitivity for better understanding of the fundamental charge generation and recombination processes in NCD films.

In the current article, the influence of the type of surface termination and boron and phosphorus doping, on the photoconductive properties of NCD films using the contactless time-resolved microwave conductivity (TRMC) technique are presented and discussed in detail. One very important characteristic of the TRMC technique is that no electrical contacts to the film are needed. As a result this approach overcomes contact problems that can be present in conventional photoconductivity measurements, including non-Ohmic contacts, contact resistance and adhesion problems. In addition to TRMC, valence band spectra of the NCD films are studied using X-ray photoelectron spectroscopy (XPS).

2. EXPERIMENTAL SECTION

Hereafter, we will refer to undoped NCD, boron-doped NCD and phosphorus-doped NCD films as undoped NCD, B-NCD and P-NCD, respectively. All the NCD (undoped and doped) films were grown on quartz substrates under similar conditions. Prior to the CVD growth, the quartz substrates were seeded with diamond nanoparticles by microwave cell and is given by\textsuperscript{25}

\begin{equation}
\Delta P(t)/P = -KΔG(t)
\end{equation}

where \(K\) represents a sensitivity factor that is determined from the dimensions of the microwave cell and the geometrical properties of the media in the microwave cell. The time-dependent change in conductance \(ΔG\) can be related to the normalized change in the reflected power \(ΔP(t)/P\) from the microwave cell and is given by\textsuperscript{25}

\begin{equation}
F_A = 1 - (I_x + I_y)/I_0
\end{equation}

where \(I_0\) is the incident light intensity.

Photoconductance measurements on the NCD films were carried out using the TRMC technique at room temperature and ambient pressure. Previously, this technique has been used to study the charge transport properties in a wide range of materials including organic blend films and (nanostructured) inorganic semiconductors.\textsuperscript{23–25} In one of our recent studies, we used the TRMC technique for the first time to study diamond films in an electrode-free way.\textsuperscript{26} A new method to hydrogenate CVD diamond films by high-temperature annealing at atmospheric pressure was introduced. To compare the resulting films with films that were hydrogenated with the established plasma-based method, we used a variety of techniques, including TRMC. A detailed description on the TRMC technique can be found in this review.\textsuperscript{25} Briefly, the home-built TRMC setup consists of an X-band (8.4 GHz) microwave cell (Figure 1). The NCD film was photoexcited with a 3 ns laser pulse from an optical parametric oscillator pumped by a Q-switched Nd:YAG laser (Vibrant II, Opotek). The overall response time of the setup including the electronics used for detection is \(\sim\)1 ns. The measurements were performed at 300 nm (4.13 eV), 580 nm (2.14 eV), and 700 nm (1.77 eV) wavelengths, which all correspond to energies smaller than the bandgap of diamond.

Photoexcitation of the NCD film results in the generation of mobile charge carriers in the film, which consequently changes the conductance \(ΔG\) of the film. Because there is no external electrode/contact to the film, the photogenerated mobile charges eventually decay with time via recombination and/or trapping. The change in conductance within the film \(ΔG(t)\) can be related to the normalized change in the reflected power \(ΔP(t)/P\) from the microwave cell and is given by\textsuperscript{25}

\[\Delta G(t) = G(t) - G(0)\]

where \(G(0)\) is the initial conductance of the film and \(G(t)\) is the conductance at time \(t\).

Figure 1. Schematic of the microwave cell used in the TRMC technique.

To estimate the average (surface) grain size and the surface roughness of the NCD films, we performed tapping-mode atomic force microscopy (AFM) with a Nanoscope Dimension 3100 scanning probe microscope using silicon cantilevers (OLYMPUS) with a resonance frequency of \(\sim\)350 kHz and a spring constant of \(\sim\)26 N/m. Raman spectroscopy was performed at 514 nm excitation wavelength with a laser spot size of \(\sim\)400 nm using a Renishaw in via micro-Raman spectrometer to identify the sp\(^3\) bonding characteristics, i.e., the diamond phase of the NCD film.

The valence band measurements of NCD films were carried out using a monochromated Al K\(_{\alpha}\) XPS (Thermo Fisher Scientific, K Alpha model). XPS measurements were performed at a base pressure of \(10^{-7}\) mbar with a spot size of 400 μm. Valence band scans were taken at 100 eV pass energy at 3 different spots on the film with each scan averaged over 10 times. The flood gun was kept on during all XPS measurements to compensate for the potential charging of surfaces. Optical transmission \((I_x)\) and reflection \((I_y)\) spectra of the NCD films were measured using a PERKIN ELMER Lambda 900 UV/vis/NIR spectrometer. The spectra were used to obtain the fraction of incident photons absorbed \((F_A)\) by the film, which is given by

\[F_A = 1 - (I_x + I_y)/I_0\]

\(I_0\) is the incident light intensity.

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\[ΔP(t)/P = -KΔG(t)\]

where \(K\) represents a sensitivity factor that is determined from the dimensions of the microwave cell and the geometrical properties of the media in the microwave cell. The time-dependent change in conductance \(ΔG(t)\) can be further expressed in terms of the number of photoinduced electrons and holes and the sum of their mobilities. If the decay of the photoinduced charges is longer than the time resolution of the detection system, then the maximum change in conductance \(ΔG_{\text{max}}\) with respect to the incident light gives the following equation

\[ΔP(t)/P = -KΔG(t)\]

where \(K\) represents a sensitivity factor that is determined from the dimensions of the microwave cell and the geometrical properties of the media in the microwave cell. The time-dependent change in conductance \(ΔG\) can be related to the normalized change in the reflected power \(ΔP(t)/P\) from the microwave cell and is given by\textsuperscript{25}
Here, $\Phi \Sigma \mu$ denotes the product of the quantum yield of the photoinduced charges ($\Phi$) per absorbed photon and the sum of electrons and holes mobilities ($\Sigma \mu$), $I_0$ is the incident light intensity per pulse, and $F_A$ is the fraction of incident photon absorbed. $\beta$ is the ratio between the broad and narrow internal dimensions of the microwave cell and $e$ is the electron charge.

3. RESULTS AND DISCUSSION

In the first part of this section, we elaborate on the film characterization, which includes WCA for surface wettability, AFM for surface topography and grain size, Raman spectroscopy for the presence of crystalline diamond phase, and XPS measurements for valence band profile. The second part focuses on the TRMC photoconductance data of hydrogen and oxygen-terminated NCD: undoped, boron-doped, and phosphorus-doped.

To evaluate the surface wetting properties, we carried out WCA measurements. For boron-doped, phosphorus-doped and undoped NCD:O films, the average static water contact angle was found to be less than 20°, whereas values of 80°−90° were found for the corresponding NCD:H films, which is in line with reported values.26

For characterizing the topography and the surface grain size of the NCD surface, we performed AFM. Figure 2 depicts AFM topography images of undoped NCD:O, B-NCD:O and P-NCD:O films showing the nanocrystalline character of the films with a high density of grains. The average surface grain sizes of the undoped NCD:O, B-NCD:O and P-NCD:O films were found to be ~60, ~128, and ~78 nm, respectively, and the root-mean-square values of the surface roughness were found to be ~17, ~29, and ~47 nm, respectively.

To identify the sp³ phase in the NCD films, we carried out Raman spectroscopy. Figure 3 presents the Raman spectra of hydrogen- and oxygen-terminated undoped NCD, B-NCD and P-NCD films with a peak (ii) at 1329, 1321, and 1330 cm⁻¹, respectively, which corresponds to the sp³ vibrational mode of diamond.27,28 The broadening and shifting of the diamond peak from the characteristic 1332 cm⁻¹ line can be associated with the grain size of the NCD film and the eventual presence of strain.27 The peaks observed at 1135 and 1475 cm⁻¹ for undoped NCD and peaks 1150 and 1477 cm⁻¹ for B-NCD (i.e., peaks i and iii in both cases) correspond to the C−H vibration modes, which also have been reported previously in these kind of films.29 The broad hump around 1550 cm⁻¹ for undoped NCD (peak iv), 1600 cm⁻¹ for B-NCD (peak iv) and 1550 cm⁻¹ for P-NCD (peak i) are attributed to sp²-bonded carbon, which is predominantly expected to be present at the grain boundaries.30,31 Finally, the peak observed at 1220 cm⁻¹ for B-NCD film (peak v) can be ascribed to boron doping.32,33

To further characterize the surface of NCD films, we carried out XPS valence band measurements. Figure 4 shows XPS valence band spectra for undoped and doped NCD films. The undoped NCD:H plot consists of 4 features as identified by ref 34 (Figure 4a, dashed lines): (i) a peak below ~11 eV is attributed to the valence bands related to the C 2p levels, which...
are linked to H 1s orbitals, (ii) a peak around \( \sim 12 \) eV that corresponds to the valence band derived from the 2s–2p hybridized levels of the diamond, (iii) a peak around \( \sim 17 \) eV related to the 2s level of the diamond and (iv) a low-intensity peak around \( \sim 25 \) eV, which is ascribed to the photoemission from O 2s.\(^{34}\) In comparison to the undoped NCD:H film, the C-related peaks (i.e., peaks i-iii) in the valence band spectrum of undoped NCD:O film are shifted toward higher binding energies. This shift can be attributed to the surface dipoles increasing the work function, which is consistent with the shift observed for C 1s core-level spectra in the undoped NCD:O film.\(^{22,34}\) In addition, the photoemission peak due to O 2s (peak iv) is higher in intensity for the undoped NCD:O film compared to the undoped NCD:H film. This difference is in agreement with the higher O 1s/C 1s atomic concentration ratio in undoped NCD:O film.\(^{22}\) Interestingly, for B-NCD:H and B-NCD:O films (Figure 4b), there is hardly any shift observed in the valence band binding energies. This could be due to the rather high boron doping in our film (boron concentration \( \sim 3000 \) ppm, bulk concentration of boron \( \approx 1.6 \times 10^{21} \) cm\(^{-3}\)), which pins the Fermi level at the boron level near the valence band.\(^{35}\) The O 2s photoemission peak in B-NCD:H is found to be slightly lower in intensity as compared to B-NCD:O, which is not significant as compared to the differences observed in undoped NCD films. However, the WCA measurements of B-NCD:H and B-NCD:O surfaces showed a clear difference in surface wettability indicating different surface termination. The valence band data of the P-NCD films is given in Figure 4c and clearly shows that the Fermi level pinning is less dominant as compared to the B-NCD films.

To conclude from the first part of the Results and Discussion, the difference in the valence band spectra of undoped NCD:H and undoped NCD:O films is in line with the difference in the related band bending profiles, due to a shift in the Fermi level. On the contrary, the valence band spectra of the B-NCD:H and B-NCD:O films were found to be very similar, which can be understood by pinning of the Fermi level at the surface.

Next, the samples were studied with TRMC. Figure 5a shows a TRMC trace obtained on the excitation of undoped NCD:H by a nanosecond laser pulse of 300 nm. From the trace two characteristic parameters can be extracted: the magnitude given by the maximum value of \( \Delta G/\beta eI_{F_A} \) indicated by the arrow and (inset) log-lin plot showing the charge carrier lifetime data (dotted line) and corresponding fit using a double exponential function (solid line). (b) Photoconductance transient of undoped NCD:O at 300 nm wavelength. The arrow indicates the maximum value of the photoconductance magnitude.

In summary, the band bending of the B-NCD:H and B-NCD:O films is increased by the boron concentration and is lower for the undoped NCD:H and NCD:O films. The photoconductance properties of NCD:H and NCD:O films are in line with the differences in work function and charge carrier lifetime, indicating the high work function and the long lifetime of the photogenerated charges due to the surface pinning by boron.
and wavelengths. Figure 6 (white section, top) shows the maximum values of the photoconductance expressed in $\Phi \Sigma \mu$ as a function of the laser pulse intensity ($I_0$) at three different sub-bandgap wavelengths, i.e., 300, 580, and 700 nm. At $I_0 > 2 \times 10^{13}$ photons/cm$^2$, the value of $\Phi \Sigma \mu$ decreases with increasing intensity. This could be due to second-order recombination reducing the number of photoinduced mobile charges or due to charge carrier scattering resulting in a lowering of the mobility.37 Interestingly, the dependence of the $\Phi \Sigma \mu$ values on $I_0$ is the same for all measured wavelengths. These results suggest that for undoped NCD:H the charge carrier generation and recombination kinetics are identical for all three photon energies.

Figure 6 (gray section, bottom) includes the maximum values of $\Phi \Sigma \mu$ for undoped NCD:O as a function of $I_0$ at the same three wavelengths. Only at intensities above $1 \times 10^{14}$ photons/cm$^2$, a small photoconductance signal could be observed. For intensities less than $1 \times 10^{14}$ photons/cm$^2$ the signal-to-noise ratio was too small to be resolved. Because an electronic transition from the valence band to the conduction band of diamond requires an energy of $\gtrsim 5.47$ eV ($\sim 226$ nm) all photon energies used in this work result in an electronic transition to or from an intrabandgap state; most likely an electron from the valence band to an empty $\pi^*$ state. The differences observed in the TRMC results obtained for undoped NCD:H and undoped NCD:O can be rationalized by a change in the band bending diagram at the interface with air, because of the differences in surface termination. It has been reported previously that the electric dipoles caused by hydrogen atoms at the surface of diamond, lead to a reduction of the surface Fermi level (toward a decreased work function). This results in an upward band bending diagram in the presence of a water film at the diamond/air interface, facilitating the dissociation of an electron–hole pair formed on photoexcitation.38–40 We postulate that upon sub-bandgap excitation of undoped NCD:H electrons are trapped in empty $\pi^*$ states leaving behind mobile holes in the valence band (Figure 7a), which are drifted toward the surface. This suggests that the obtained mobility value of $\sim 0.27$ cm$^2/(V \cdot s)$ from the TRMC measurements is due to mobile holes, comparable to the hole mobility reported for both boron-doped and undoped hydrogenated NCD films.8,21,36 The fact that the generation yield is independent of the three, largely varying wavelengths suggests that photoexcited electrons are trapped in common empty $\pi^*$ states. In the case of undoped NCD:O, oxygen termination results in a downward band bending diagram due to the opposite electric dipole moment of oxygen as compared to hydrogen termination (Figure 7b).40,41 Photoexcited electrons residing in $\pi^*$ states are immobile, while the corresponding holes remain in the bulk of the grains of the NCD. Hence, fast second-order electron–hole recombination can take place explaining the rapid TRMC decay of $\sim 100$ ps. The band-bending diagram presented here to explain the TRMC results agrees well with the XPS valence band measurements on undoped NCD:H and undoped NCD:O (Figure 4a) discussed above, which show a shift in the surface Fermi levels.

To study the effect of doping on the optoelectronic properties of NCD, we performed TRMC measurements on boron (B-NCD) and phosphorus (P-NCD) doped films. Figure 8a shows the TRMC traces of hydrogen (B-NCD:H) and oxygen (B-NCD:O) terminated NCD, which interestingly are found to be similar. If $\Phi$ is assumed to be 1, the lower limit of the mobility is $\sim 0.05$ cm$^2/(V \cdot s)$. Note that although a high boron concentration is used (boron concentration $\sim 3000$ ppm, bulk concentration of boron $1.6 \times 10^{21}$ cm$^{-3}$) the majority of...
undoped and doped NCD films. The photoconductance signal and the charge carrier lifetimes are significantly affected by the type of surface termination on the undoped NCD films, whereas no surface termination effect was observed for boron-doped films. These results are explained on the basis of the differences in the band bending diagrams at the diamond surfaces. The photoconductance results from the TRMC technique provide a direct quantification of the charge carrier mobility and its decay dynamics without problems related to electrode contact issues. Furthermore, the charge transport study presented here provides valuable information to investigate the effect of surface functionalization on NCD films for potential surface-based (opto)electronic sensor applications and the charge transfer between diamond-organic systems for solar cell applications.35

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Figure 8. Photoconductance transients of (a) B-NCD and (b) P-NCD at 300 nm wavelength. Note the different time scales. The inset in (a) shows the charge carrier lifetime data for the B-NCD film.


