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Characteristics of zirconium and niobium contacts on boron-doped diamond

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Abstract

Formation of ohmic contacts on boron-doped diamond using zirconium and niobium has been studied in comparison to the commonly used titanium or tantalum contacts. Metal contacts were fabricated using standard micro-fabrication technologies on epitaxial layers with different boron concentrations. Room temperature specific contact resistance was determined using the circular Transmission Line Model after annealing at various temperatures. The specific contact resistance varies considerably with boron concentration and annealing temperature. Zirconium and niobium form ohmic contacts on highly boron-doped diamond after high-temperature annealing with a specific contact resistance comparable to that of titanium and tantalum contacts. Furthermore, we observed zirconium contacts showed better thermal stability up to 700 °C upon high-temperature annealing compared to titanium and other metal contacts. Finally, these results emphasize the potential of zirconium for the formation of ohmic contacts on boron-doped diamond.

Keywords: Ohmic contact, boron-doped diamond, circular transmission line model, zirconium, niobium, titanium

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1 Introduction

Diamond is a semiconductor with unique properties [1], which has been foreseen as an advantageous replacement for silicon in the fabrication of high frequency and high power devices such as high voltage Schottky diodes [2,3] and more recently MOSFETs [4] and other electronic devices [5–8]. Stable ohmic contacts with a low specific contact resistance are crucial to limit electric losses in the operation of electronic power devices. Numerous factors, such as the nature of the metal [9–16], the doping level [12], the surface pre-treatment [9,17–19], the crystalline orientation [16], and the annealing time and temperature [12,20–22] are important in the formation of ohmic or Schottky contacts. The ohmic behavior of the contacts is generally associated related to carbide formation [11–13,23–25]. A large variety of carbide forming metals have already been studied for the fabrication of ohmic contacts on boron (B) doped diamond [10,11,26–30]. Titanium (Ti) is generally preferred for this purpose.

With the aim of studying impurity impact ionization and avalanche effects in B-doped diamond in high electric fields [31,32], we searched carbide forming refractory metals with a high chemical affinity to carbon, similar to titanium, capable of forming good ohmic contacts,
which could withstand high current densities and high temperatures. Niobium (Nb), tantalum (Ta) and zirconium (Zr) are refractory metals with a Gibbs free energy for carbide formation close to that of Ti [33,34]. While Ta has been already studied, the formation of ohmic contacts on B-doped diamond using Nb or Zr has not, according to our knowledge, been reported in the literature. On the contrary, Zr has been reported to form Schottky contacts [2,35]. In this work, we studied the formation of ohmic contacts on B-doped diamond using Zr and Nb metals in comparison to the well-studied Ti and Ta metals. Epitaxial B-doped diamond layers were grown by microwave plasma enhanced chemical vapor deposition with various B concentrations. Metal contacts were patterned by evaporation and standard lithography processes. The current-voltage (I-V) characteristics of different metal contacts were measured at room temperature, after annealing at various temperatures, and for different B concentrations. The diamond layers’ resistivity, contact resistance, and specific contact resistance were determined using the circular Transmission Line Model (c-TLM) [36,37].

2 Material and methods

B-doped epitaxial diamond layers were grown in a microwave plasma enhanced chemical vapor deposition (MW PECVD) AX5010 reactor using a methane, hydrogen and trimethylborane gas mixture. The boron to carbon ratio (B/C ratio) in the gas phase has been varied from 500 ppm to 20,000 ppm to obtain B-doped epitaxial diamond layers between a few $10^{19}$ cm$^{-3}$ to a few $10^{21}$ cm$^{-3}$, i.e. from the semiconductor conduction region below metallic transition up to highly doped diamond with metallic conduction. B-doped epitaxial diamond layers, with B/C ratios of up to 6000 ppm, were grown on 3×3×0.8 mm$^3$ (100)-oriented high-pressure high-temperature (HPHT) grown synthetic single crystal diamond substrates using the process reported in Ref. [38]. Layers grown using this process exhibit nearly defect-free growth with a very low roughness of 0.2 nm and 1 nm over areas of 5×5 μm$^2$ and 90×90 μm$^2$, respectively (see Figure 1). Above the threshold of B/C 6000 ppm, epitaxial diamond growth is no longer sustained on (100)-oriented substrates. Hence B-doped epitaxial diamond layers with B/C = 20,000 ppm were grown on 2×2×0.8 mm$^3$ (111)-oriented HPHT grown synthetic single crystal diamond substrates using deposition conditions reported in Table 1. In contrary to B-doped epitaxial diamond layers grown on (100)-oriented substrates, layers grown on (111)-oriented substrates were not etched in an oxygen-hydrogen plasma before diamond deposition. Prior to Before and after MW PECVD deposition, samples were cleaned in a hot admixture of sulfuric acid and potassium nitrate for 10 min. The samples were then rinsed with hot deionized water, acetone and isopropyl alcohol in an ultrasonic bath for 10 min and finally dried with clean compressed air. Several layers were grown with the same B/C ratio to enable the study of the properties of contacts fabricated with the four different metals. Thicknesses of the B-doped epitaxial diamond layers, used to compute the layer resistivity, was determined from double mass measurement using a high precision ME5 microbalance (Sartorius).

Circular Transmission Line Model test structures were fabricated by photolithography and metal evaporation. c-TLM structures were preferred as they are easily realized without the need of a MESA etch to prevent current spreading in linear TLM structures. The electrodes were patterned by the lift-off technique using a positive photoresist ma-P 1210 from Microresist technology exposed with a mask-less laser lithography system: Microwriter ML
from the Lot Oriel company. Details of the lithography process are described in ref. [39]. The spacing between the inner circular electrodes (150 μm in diameter) and the outer electrodes varies between 10 and 60 μm (see Figure 1). Metals were deposited by e-beam evaporation in an Edwards Auto 500 vacuum coater. Zr, Nb, Ta and Ti 10 nm thick contacts were capped using a 90 nm thick gold layer with a 10 nm thick platinum inter-diffusion layer. The I-V characteristics of fabricated c-TLM structures were characterized using a computer controlled Agilent 4142B modular DC source/monitor and a probe station Suss Microtech PA-200. Electrical characteristics were measured at room temperature for as-deposited and consecutively annealed contacts with increasing temperature: 200 °C, 400 °C, 500 °C, 600 °C and 700 °C. Samples were annealed at low pressure in a rapid thermal annealing system, AS-One from Annealsys for 20 min in a flow of nitrogen.

The resistance of c-TLM structures has been calculated using linear least squares fitting I-V characteristics. Due to the circular geometry, the non-linear variation of the resistance as a function of the gap spacing is linearized using a correction factor (see Figure 2). The correction factor (c) is a function of the inner electrode radius (R) and the inter-electrode spacing (s) as described in equation (1).

\[ c = \frac{R}{s} \ln \left( \frac{R+s}{R} \right) \]  

Linear least squares fitting of corrected resistance enables the determination of the contact resistance \( R_c \), transfer length \( L_T \), and the diamond sheet resistance \( R_{sh} \) from the Y-axis intercept, the X-axis intercept and the slope, respectively [36].

### 3 Results and discussion

The resistivity of B-doped epitaxial diamond layers extracted from c-TLM as a function of the B/C ratio is reported in Fig. 3. Based on the ref. [40,41], one can cautiously assume conservation of the B/C ratio from the gas phase to the solid phase during deposition to estimate boron incorporation into grown layers. This assumption is supported by the quantitative agreement of measured values with values from the literature [42,43]. The resistivity drops from a few \( \Omega \cdot \text{cm} \) in the region of incomplete ionization, i.e. when B/C = 500 ppm, to a few 10s of m\( \Omega \cdot \text{cm} \) for B/C ≥ 2000 ppm in the region of metallic conduction. The higher resistivity values of epitaxial layers prepared with B/C = 20,000 ppm are attributed to the known poorer conduction properties of B-doped epitaxial diamond layers grown on (111)-oriented substrates [40]. As deposited Ta, Nb and Zr contacts fabricated on B-doped epitaxial diamond layers grown with a B/C ratio of 500 ppm exhibit symmetric but non-ohmic I-V characteristics, which is in contrary to Ti contacts (not shown). Nb and Zr form ohmic contacts only after high temperature annealing (≥ 400°C). The evolution of I-V characteristics with annealing temperature of layers grown at high B concentration (B/C = 20,000 ppm) is shown in Fig. 4. Remarkably, as deposited Ta also formed ohmic contact on highly doped layers (B/C = 20,000 ppm) with a ca. 2×10^4 \( \Omega \cdot \text{cm}^2 \) specific contact resistance. The total resistance of Ti contacts suddenly increases after high temperature annealing (≥ 600 °C). This increase in resistance indicates a degradation of the specific contact resistance. The variations of the calculated specific contact resistance as a function of the annealing temperature and the B concentration for the different fabricated metal contacts are reported in Fig. 5 and Fig. 6. Fig. 6 shows that a clear decrease in the specific contact resistance with the rising B concentration is observed for all metals. The specific contact resistance drops by almost four orders of magnitude from a few
Ω.cm² to a few 10⁻⁴ Ω.cm² and below as the B concentration increases by a factor of 40 from 500 ppm to 20,000 ppm. The lowest contact resistance of 4×10⁻⁵ Ω.cm² was obtained for an as-deposited Ti contact on a layer grown with B/C = 20,000 ppm. Note that all contacts on heavily B-doped epitaxial diamond layers, i.e. above the metallic transition [44], exhibited low specific contact resistance values of about 10⁻⁴ Ω.cm² after optimal annealing. The same decrease in specific contact resistance of Ti and Mo contacts on polycrystalline B-doped diamond was reported by Nakanishi et al. who also reported a linear correlation between specific contact resistance and the resistivity of the doped layer [22]. The strong influence of acceptor concentration (N_a) is associated to tunneling transport at the contact interface [22] and the N_a⁻¹/² decrease of the Schottky contact depletion width [45]. The specific contact resistance of Ti and Ta contacts decreases steadily with annealing temperature between 200°C and 500°C. However, B-doped epitaxial diamond layers grown at a high B/C ratio, e.g. of 20,000 ppm, show that a rapid increase in the specific contact resistance is observed after annealing at higher temperatures, which indicates a degradation of the ohmic contacts. This effect is also observed in Nb contacts. Similar degradation of ohmic contacts observed for Ti and Mo contacts on B-doped diamond upon annealing was reported by Johnston et al. [14]. Whilst less pronounced, degradation of Ti, Ta and Nb contacts is also observed on layers with lower B concentrations. In contrary to this, the Zr metal-zirconium contacts show a steady improvement of ρ_c values with increasing temperature up to 700°C. On the contrary, Zr metal-zirconium contacts show a steady improvement of ρ_c values with increasing temperature up to 700°C. The higher thermal stability of Zr contacts suggests that they might have a longer lifetime before failure than conventionally used Ti contacts. The optimal specific resistance values of Ti, Nb, Zr, and Ta contacts for the various B concentrations studied in this work and their respective annealing temperature are summarized in Table 2. These results clearly show that Nb and Zr metals form ohmic contacts with specific contact resistance comparable to the ordinarily used Ti or Ta metals. Specific resistances reported in this work are larger than reported values from the literature [22]. This difference is attributed to the absence of an amorphisation process step in the preparation of the metal contacts [16,19,46,47].

**Conclusion**

The properties of Ti, Ta, Zr and Nb metal contacts on B-doped epitaxial diamond layers have been measured and compared using circular transmission line model for various B concentrations and different consecutive annealing temperatures. Doped diamond layer resistivity values determined from c-TLM are consistent with data from the literature. Formation of ohmic contact using Zr and Nb has been demonstrated with specific contact resistance comparable to the ones of Ti and Ta after high-temperature annealing. The specific contact resistance varies significantly with B concentration, nature of the metal contact, and annealing temperature. Finally, we observed Zr contacts on highly boron doped diamond exhibit better stability upon high-temperature annealing compared to other studied metallic contacts, especially titanium contacts. This result suggests that Zr contacts might advantageously replace conventional Ti contacts for a long lifetime and high-temperature applications of diamond devices.

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References


Figure 1: Scanning electron microscopy images of (a) B-doped epitaxial diamond layer on (100)-oriented HPHT substrate and (b) c-TLM structures fabricated on a B-doped epitaxial diamond layer deposited on a (111)-oriented HPHT substrate. 90×90 μm² AFM topography image of a B-doped epitaxial diamond layer grown on a (100)-oriented substrate (c).

Table 1: Diamond deposition conditions

<table>
<thead>
<tr>
<th>Substrate orientation</th>
<th>MW power (W)</th>
<th>Pressure (mbar)</th>
<th>CH₄ in H₂ concentration</th>
<th>Deposition temperature (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>(100)</td>
<td>700</td>
<td>100</td>
<td>1%</td>
<td>1100</td>
</tr>
<tr>
<td>(111)</td>
<td>575</td>
<td>60</td>
<td>0.1%</td>
<td>1000</td>
</tr>
</tbody>
</table>
Figure 2: Total resistance vs electrodes’ gap spacing before (squares) and after (red circles) correction for the circular geometry.

Figure 3: Room-temperature resistivity of (100) and (111) B-doped epitaxial diamond layers as a function of the boron to carbon ratio B concentration in the gas phase with respect to the carbon source. Comparison between literature (red dash line) [42,43] and experimental data from this study (the boron concentration ([B]) of the upper x-axis is calculated was determined assuming conservation of B/C ratio from the gas phase to the solid phase during diamond deposition [41,48]).
Figure 4: I-V characteristic of Ti, Nb, Zr and Ta contacts on B-doped diamond annealed at several temperatures up to 700°C.
Figure 5: Specific contact resistance as a function of annealing temperature for Ti, Nb, Zr, and Ta contacts to various B-doped epitaxial diamond layers.

Table 2 Optimal specific contact resistance of the Ti, Nb, Zr and Ta contacts for various B concentrations after annealing at optimal temperatures.

<table>
<thead>
<tr>
<th>B/C vs $\rho_c$</th>
<th>Ti ($\Omega$.cm$^2$)</th>
<th>Nb ($\Omega$.cm$^2$)</th>
<th>Zr($\Omega$.cm$^2$)</th>
<th>Ta($\Omega$.cm$^2$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>500 ppm</td>
<td>1.4 (600°C)</td>
<td>1.1 (500°C)</td>
<td>0.3 (700°C)</td>
<td>5.9 (700°C)</td>
</tr>
<tr>
<td>2,000 ppm</td>
<td>8.5×10$^{-4}$ (200°C)</td>
<td>9.7×10$^{-2}$ (500°C)</td>
<td>0.2 (400°C)</td>
<td>1.3×10$^{-2}$ (400°C)</td>
</tr>
<tr>
<td>6,000 ppm</td>
<td>1.7×10$^{-4}$ (25°C)</td>
<td>1.4×10$^{-4}$ (25°C)</td>
<td>1.2×10$^{-2}$ (500°C)</td>
<td>12.7×10$^{-3}$ (700°C)</td>
</tr>
<tr>
<td>20,000 ppm</td>
<td>4.1×10$^{-5}$ (500°C)</td>
<td>2.5×10$^{-4}$ (600°C)</td>
<td>5.5×10$^{-4}$ (700°C)</td>
<td>1.5×10$^{-4}$ (400°C)</td>
</tr>
</tbody>
</table>
Figure 6: Specific contact resistances of Ti, Nb, Zr, and Ta contacts annealed at different temperatures as a function of B/C ratio during deposition and boron concentration [B] (the boron concentration ([B]) of upper x-axis is calculated assuming conservation of B/C ratio from the gas phase to the solid phase during diamond deposition [41,48]).