Multiple conduction paths in boron δ-doped diamond structures

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Impedance spectroscopy has been used to investigate conductivity within boron-doped diamond in an intrinsic/delta-doped/intrinsic (i-δ-i) multilayer structure. For a 5 nm thick delta layer, three conduction pathways are observed, which can be assigned to transport within the delta layer and to two differing conduction paths in the i-layers adjoining the delta layer. For transport in the i-layers, thermal trapping/detrapping processes can be observed, and only at the highest temperature investigated (673 K) can transport due to a single conduction process be seen. Impedance spectroscopy is an ideal nondestructive tool for investigating the electrical characteristics of complex diamond structures. © 2009 American Institute of Physics. [DOI: 10.1063/1.3075860]

Diamond is considered to be an ideal material for the fabrication of high performance electronic devices if problems associated with poor material quality and doping can be overcome.¹ Recent single crystal diamond layers grown by plasma-enhanced chemical vapor deposition (CVD) have been shown to display carrier mobility values that surpass those of the very best natural diamond stones.² Doping remains a more significant problem; boron forms a p-type acceptor with an activation energy (Ea) of 0.37 eV, meaning that few holes are present at room temperature.³ Borst and Weiss⁴ showed that when [B] > 10¹⁸ cm⁻³, this energy sharply declines, as a boron impurity band emerges, with concentrations greater than 10¹⁹ cm⁻³, leading to semimetallic characteristics. While this removes the problem associated with the Ea of the acceptors, the conduction mechanism has now become “hopping” with the associated extremely low values of carrier mobility. A solution may be the use of δ-doping, where a highly boron-doped layer with nanometer dimensions is sandwiched between undoped materials. The heavily doped δ-region will provide carriers with little thermal activation, while the carriers will emerge from this layer and be transported through the (higher mobility) intrinsic layers each side. A number of reports of the growth of such structures and their use for the formation of field-effect transistors (FETs) have been made and this work has been reviewed.⁵,⁶ The ideal δ-layer thickness would be 1–2 nm.⁷ Given the nature of the CVD process used this is technologically challenging. Aleksov et al.⁷ reported a thickness of 6 nm, although recently the same group achieved 1.1 nm,⁸ corresponding to around three atom layers of doped diamond.

It is essential to be able to determine the electronic characteristics of the δ-layer and the surrounding intrinsic material if effective FETs are to be designed. Hall effect measurements can provide carrier concentrations and mobility values, but understanding the contribution of each region to the numbers determined this way is difficult. Aleksov et al.⁷ reported a method whereby repeated etching-measurement cycles were used to study the an i-δ-i structure, revealing an activation energy of 30 meV within the δ and 1.5 eV outside. However, this approach is time-consuming, limited in depth resolution by the etching process, and destructive of the layers. In this letter we report on the use of impedance spectroscopy (IS) for determining conduction paths within complex diamond i-δ-i structures and their characteristics.

Microwave plasma-enhanced CVD was used to produce both intrinsic and boron-doped layers on type Ib high pressure high temperature single crystal substrates using growth conditions for high quality materials similar to those described previously.⁷ Following the growth of a 100 nm thick i-layer on the substrate, a δ-layer was grown with a full width at half maximum thickness of 5 nm as judged by secondary ion mass spectroscopy (SIMS). Onto the δ-layer was grown an intrinsic capping layer of 20 nm thickness. SIMS analysis showed [B] in the δ-layer to be ~5 × 10²⁰ atoms cm⁻³. All samples were subjected to treatments that are known to leave the surface in a strongly oxidized state,⁹ avoiding so-called “surface conductivity.”¹⁰ Au contacts were evaporated onto the surface of the samples, which were then placed inside a vacuum chamber providing atmospheric control and electrical shielding during analysis. Measurements were made within the temperature range 300–900 K. IS was performed with a Solartron SI1260 with a high impedance input module 1296. The real and imaginary parts of the impedance were measured as a function of frequency (0.1 Hz–10 MHz) to produce “Cole–Cole” plots. Each contribution to the overall impedance of the sample can be characterized by an individual RC component and hence differing semicircular responses in the Cole–Cole plots. In practice these responses often overlay each other, meaning that careful fitting of the data to ideal semicircles is required.¹¹ The measurements are repeated as a function of temperature such that the activation energy for each conduction path can be determined.

Cole–Cole plots for a typical sample are shown in Figs. 1(a)–1(c) for increasing sample temperature (not all temperatures are included for brevity). In Fig. 1(a) the imaginary component of the impedance is plotted against the real component as a function of frequency at 300 K. Initial inspection
of the graph suggests the presence of a single semicircular response, while in Fig. 1(b) the situation is clearly somewhat more complicated once the temperature has increased. First, the inspection of the values on the axes shows that the film has become more resistive as the temperature increases from 300 to 373 K. Further, the shape of the response has altered. Moreover, impedance measurements for a temperature of 423 K show a further increase in resistance values and a rather dramatic change in the shape of the graph [Fig. 1(b)]. Also shown in Fig. 1(b) are the measurements obtained for a temperature of 473 K. Intriguingly, the impedance values are now lower than those for 373 and 423 K, so the sample now becomes less resistive as the temperature further increases. This trend continues as the temperature is raised to 573 K [Fig. 1(c)] with little apparent change at 673 K. To confirm that these observations did not relate to the type Ib substrate underlying the i-B-i structure, impedance measurements were obtained for a Ib crystal alone; the resistance remained considerably greater than that recorded here (at greater than $10^9$ Ω) over this temperature range.

To model the $RC$ components of each conduction path leading to the data shown in Fig. 1, it is necessary to fit semicircular responses to the data. A semicircular response can be fitted to the data before the curve rises steeply in value as the frequency decreases. A second semicircular response can then be fitted, followed by a third as the frequency is decreased further. Table I shows the $R$ and $C$ values that can be determined from these fits. In fact, while the magnitude and relative contributions vary with temperature, a similar analysis of all temperatures (including 300 K) up to 523 K showed the clear presence of three semicircular responses in all of the impedance measurements. However, the data presented in Fig. 1(c) for analysis at 673 K revealed only one semicircular response.

It is interesting to consider the way the resistance determined for each of these $RC$ contributions changes with temperature such that an activation energy for each process can be determined. Figure 2 shows this data with the resistance of each plotted against $1/T$. It is immediately apparent that semicircle 1 has the lowest resistance and displays little variation in $R$ with $T$; an $E_a$ of less than 20 meV can be proposed. In stark contrast the data points for both semicircular responses 2 and 3 show an increase in resistance with increasing temperature until peaking around 423 K, after which the resistance is seen to decline with increasing temperature. In the case of semicircular response 2, sufficient

![FIG. 1. Cole–Cole plots of real vs imaginary component of the impedance values determined as a function of frequency for measurements carried out at (a) 300 K, (b) 373–473 K, and (c) 573–673 K.](image)

![FIG. 2. Plot of resistance vs $1000/T$ to reveal the activation energy for the three conduction processes derived from temperature dependent Cole–Cole plots.](image)

### Table I. Resistance and capacitance values determined for the three semicircular responses within the Cole–Cole plots measured at 373 K using curve fitting.

<table>
<thead>
<tr>
<th>$T$ (K)</th>
<th>Impedance</th>
<th>Semicircle 1</th>
<th>Semicircle 2</th>
<th>Semicircle 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>373</td>
<td>$R$ (Ω)</td>
<td>13 223</td>
<td>9.7 $\times$ 10^6</td>
<td>8.6 $\times$ 10^7</td>
</tr>
<tr>
<td>373</td>
<td>$C$ (F)</td>
<td>2 $\times$ 10^{-12}</td>
<td>3 $\times$ 10^{-11}</td>
<td>3 $\times$ 10^{-10}</td>
</tr>
</tbody>
</table>
data points exist to determine that the activation energy for
the initial increase in resistance with temperature is 0.25 eV,
while the activation energy for the decreasing resistance as
the temperature increases further is 1.75 eV.

IS has been previously used to characterize polycrystal-
line diamond films,12,13 where the contributions from the
grains and grain boundaries were observed. The strength of
the impedance technique lies in its ability to identify the
individual components that contribute to the overall conduc-
tivity; we have previously shown that this can include differ-
ent contributions to conduction within doped single crystal
diamond,14,15 but the complexity of the current samples is
unprecedented. The observation of three semicircular re-
sonances up to 523 K is interesting and suggests that three
distinct conduction pathways exist within this structure. It
has been widely established that capacitance values in the pico-
farad range arise from ordered material, while contributions
from disordered material are most normally observed to lie
in the nanofarad region.11 In the present case capacitance
values for the highest frequency component are observed in
the picofarad range, while the second and third semicircular
components appear at higher capacitance values, 10 and 100
picofarad, respectively (Table I). This might suggest an in-
crease in crystal disorder for these conduction paths. The
contribution to the overall conduction that shows an activa-
tion energy of less than 20 meV is most likely to arise from
the heavily boron-doped δ-layer, as this value is typical for
doping levels in the region of 1020 cm−3 as we have here.4
That the resistance of this region is so low also agrees with
this assignment. The other conduction paths that can be ob-
erved are considerably more resistive, implying that the car-
rier densities responsible are comparably low. That both of
the other conduction paths display two activation energies
with opposite signs is intriguing. To increase the resistance
of the conduction path with increasing temperature implies
that the carriers responsible are acquiring sufficient energy to
become trapped at sites with an energy of some 0.25 eV from
the valance band edge, assuming the process to be one of
hole transport. At temperatures greater than 423 K, this trap
state is emptied and the resistance begins to rapidly fall with
increasing temperature. The thermal activation energy for the
decrease is 1.75 eV. It is beyond the remit of this short letter
to begin to assign the states that may give rise to these en-
ergies, but a logical proposal based upon the i-δ-i structure
being investigated would be to propose that they arise from
carrier transport within the (defective) interfacial regions be-
tween the intrinsic material and the heavily doped diamond
layer. That two distinct conduction pathways can be deter-
mined would be most easily explained if transport in the
interface between the i-layer under the heavily doped dia-
mond differed from that in the capping i-layer above it. This
might be anticipated as epitaxial growth of i-diamond onto
such a heavily doped layer could easily lead to an interface
that differs from that achieved when a heavily doped boron
layer is grown on an already present high quality diamond
region. This could also explain the fact that these two con-
duction paths differ in capacitance value by an order of mag-
nitude (Table I), which may arise if one has a higher level of

crystal disorder than the other. In the current case, at 673 K
only one semicircular response can be seen [Fig. 1(e)]. The

capacitance value for this single semicircle is in the pico-
farad range, and the resistance is very low. These facts sug-

gest that at this temperature only a single conduction path is
active. This may be because all interfacial traps are emptied
at this high temperature and carriers are free to move within
a single channel. It is not possible from this data alone to
suggest whether carrier transport at this point is solely within
the delta-layer or the adjoining (now uncharged) i-layers.

In summary, we have demonstrated that IS is a powerful
tool for investigating conduction paths within complex
multilayer structures fabricated from diamond, such as the
i-δ-i structure studied here. It has been shown that the tech-
nique is sensitive to differences in the transport characteristics
displayed by the two different interfacial regions, in ad-
inction to conduction within the δ-layer itself. It is clear that
the use of IS, when allied to other characterization tools, can
provide an excellent, nondestructive method for optimizing
these types of complex diamond structures.

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