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FAST TRACK COMMUNICATION

Generation of microdischarges in diamond substrates

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Abstract
We report the generation of microdischarges in devices composed of microcrystalline diamond. Discharges were generated in device structures with microhollow cathode discharge geometries. One structure consisted of an insulating diamond wafer coated with boron-doped diamond layers on both sides. A second structure consisted of an insulating diamond wafer coated with metal layers on both sides. In each case, a single sub-millimetre hole was machined through the conductor–insulator–conductor structure. The discharges were generated in a helium atmosphere. Breakdown voltages were around 500 V and discharge currents in the range 0.1–2.5 mA were maintained by a sustaining dc voltage of 300 V.

(Some figures may appear in colour only in the online journal)

1. Introduction

Research into microdischarges, defined as discharges in which at least one dimension is less than 1 mm, has expanded considerably in recent years and the last decade has seen these devices developed for many different applications [1, 2]. Recent research has seen the investigation of different operating modes, with some authors focusing on device scaling, especially ignition processes [3, 4], while others have investigated operating regimes associated with reproducible instabilities [5–8]. Much of this research has focused on the gas-phase processes that occur in the micro-volumes inside these devices but the role of the surface processes and plasma–surface interactions is gaining increasing attention [9].

Most microdischarge devices reported in the literature consist of metal electrodes separated by an insulating ceramic such as aluminium oxide. A notable exception is the work of Eden et al [10, 11], who developed microdischarge arrays based on silicon, produced using technology developed for microelectronics and integrated circuit fabrication. Full understanding of microdischarge physics, including ignition and instability behaviour, requires operation of devices consisting of a wider range of materials. This is particularly important for small dimension devices, in which the surface area to volume ratio becomes correspondingly large. Although there has been some effort to investigate material influences in microscale devices that operate in pre-breakdown non-self-sustaining modes [12, 13], most development of stable glow-like microdischarges has focused only on a small range of materials.

The thermal, electrical and mechanical properties of diamond make it an attractive material for fabrication of micrometre-size discharge devices. First, the high thermal conductivity of diamond should enable the high gas temperatures generated in microdischarges to be effectively conducted away from the surfaces in contact with the plasma. Second, diamond in its undoped form is an excellent insulator, while doped forms of diamond can be used as electrically conducting layers. This opens up the possibility of all-diamond structures that can be fabricated as a single integrated structure, with the additional advantage that the different layers have well-matched thermal expansion. Third, the hardness and relative inertness of diamond surfaces may enable development of devices that are resistant to
thermal and electrical degradation by prolonged exposure to microdischarge plasmas. This could lead to devices with extremely long lifetimes. Finally, the properties of the diamond surfaces, such as the secondary electron emission yield, can be varied by changing the fabrication process [14, 15].

In this communication, we report successful fabrication and operation of microhollow cathode discharge devices based on microcrystalline diamond. Discharges were generated in two devices with similar geometries but different material combinations. In one device, metal electrodes were deposited onto the faces of an undoped microcrystalline diamond wafer. In a second device, the metal electrodes were replaced by layers of p-type microcrystalline diamond, heavily doped with boron, deposited by chemical vapour deposition (CVD).

Figure 1 shows the structure and geometry of the devices. Each device is based on an undoped mechanical-grade polycrystalline CVD diamond freestanding wafer purchased from Element Six (product code: 145-500-0015), polished on one side. The wafer dimensions are 250 $\mu$m x 10 mm x 10 mm. This was cleaned in warm concentrated nitric acid to remove any residues remaining on the surface after the polishing process.

The metal–diamond–metal device was fabricated using the following process. The wafer was first subjected to a dehydration bake of 30 min on a hot plate, after which layers of titanium (100 nm) and gold (500 nm) were deposited in an SVS E-Gun evaporator. The edges of the wafer were protected with kapton adhesive tape so that a non-conducting border remained between the metal coating and the wafer edge. After coating, the metal layers were annealed using a rapid thermal annealing oven (500 °C for 2 minutes). A circular hole with 200 $\mu$m diameter was then drilled through the centre of the wafer using a micro-machining system (Micronanics Ltd) based on a 355 nm diode-pumped solid-state laser. Finally, the sample was cleaned in an oxygen plasma (PVA TePla America Inc.) for 60 minutes.

Figure 2. Magnified images of the machined holes for the cases of (a) the device with metal electrodes, showing the 200 $\mu$m hole, and (b) the device with conducting diamond electrodes, showing the 300 $\mu$m hole.

A Nd : YAG laser-based micro-machining system (Alpha, Oxford Lasers, UK) was used to drill a 300 $\mu$m-diameter hole through the centre of the sample to produce the microplasma cavity. 1 mm was also laser trimmed from each edge of the sample to ensure there was no electrical short-circuit between the upper and lower B-doped layers from unintentional B contamination of the edges. Lastly, the sample was cleaned in hot (≈80 °C) 90% sulfuric acid + potassium nitrate (2 g in 40 ml of acid) for 15 minutes. This removed any contamination along with any surface graphitization resulting from the laser etching, and left the diamond surfaces oxygen terminated.

Figure 3 shows images of the two devices, obtained using a digital microscope. It can be seen that although the metal layer is slightly discoloured at the edges of the hole, the laser machining process has produced neat circular apertures for both devices.

For each device, plasma ignition was investigated using a chamber back-filled with high-purity helium gas after evacuation to <1 Pa. Electrical contact was achieved by mechanically pressing tantalum foil strips to each surface. These foil strips were, in turn, connected to the output of a dc high-voltage power supply outside the chamber. A 136 kΩ ballast resistor was placed in series with one electrode. Electrical measurements were made using high-voltage probes while photographic images were obtained using a digital microscope.

Figure 3 shows images of the plasma generated using the device with metal electrodes, obtained when the gas pressure was 10 kPa (≈75 Torr). The two images show the plasma for different values of discharge current. Figure 3(a) shows the discharge in a very low-current mode with the plasma confined to the centre of the hole, while figure 3(b) shows the plasma generated for a much higher current, where the plasma has expanded outside the hole over the electrode surface. In both images, the anode side of the discharge is shown.
Figure 3. Images of discharges generated by the device with metal electrodes for case of (a) $V = 420\,\text{V}$, $I = 0.02\,\text{mA}$ and (b) $V = 590\,\text{V}$, $I = 0.60\,\text{mA}$. The background gas was helium at a pressure of 10 kPa.

Figure 4. (a) Anode view and (b) cathode view of a discharge generated by the device with semiconducting diamond electrodes, for the case of $V = 410\,\text{V}$, $I = 0.40\,\text{mA}$. The background gas was helium at a pressure of 10 kPa.

Figure 5. $I$–$V$ curves obtained at different pressures for (a) the device with metal electrodes and (b) the device with conducting diamond electrodes. Each curve was obtained for increasing applied voltage.

impedance compared with that of the ballast resistor and the discharge itself. For the all-diamond device, however, the electrical contacts to each diamond surface are metal–semiconductor ones and the impedance of these cannot be neglected [17]. Additionally, there will be resistance between the contact and the aperture where the discharge is located. Both the nature of the contact and the conductivity of the conducting diamond layer depend on temperature and hence are difficult to estimate in a simple way. These features will be studied and characterized in a subsequent study. It is clear, however, that the $I$–$V$ curves shown in figure 5 are consistent with those observed in other studies and indicate that stable, reproducible microdischarges can be generated in these diamond-based structures.

Finally, although we have operated these devices for a relatively short period of time, we can state that each has been operated for more than 5 h without failure. This relatively long lifetime indicates that we can expect these and similar devices to be suitable for more detailed studies of the influence on discharge properties of different material surfaces. It is also promising for potential applications.

The successful operation of diamond-based micro-discharge devices represents a new technology for
Although it was not possible to draw firm conclusions about the effect of the diamond material on the characteristics of the observed microdischarges, it is still possible to conclude that both devices generate microdischarges typical of those observed in microhollow cathode discharges with similar geometry. This demonstration of microdischarges in diamond-based devices offers the opportunity for detailed investigation of the role of material properties in the behaviour of microdischarges. The new technology also has the potential to enable development of a new generation of long-lifetime microdischarge devices.

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