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Single Crystal Diamond Schottky Photodiode

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1. Introduction

Thanks to its extreme optical and electronic properties, diamond appears to be a promising semiconducting material for photon detection. Its wide band-gap, 5.5 eV, results in a very low leakage current and its electronic properties as high carrier mobility allow fast time response (J. E. Field, 1979). Besides, it has a large breakdown electric field (∼10 V/μm), a low dielectric constant (i.e. low capacitance), chemical inertness and low intrinsic carrier density, which makes cooling for noise reduction unnecessary (J. Prins, 1997). Its extreme radiation hardness is well known and another interesting feature, again related to the wide band-gap, is its selective sensitivity to radiation with wavelengths shorter than 225 nm (visible-blind detectors) (J. F. Hochdez et al., 2002). Several attempts have been made to build up UV detectors from natural or synthetic diamonds grown by Chemical Vapour Deposition (CVD). A detector often reported in literature is the photoresistor (A. Balducci et al., 2005; T. Teraji et al., 2004) having a planar structure and consisting of a photoconductive diamond film with metal electrodes placed on the top surface. It can operate only with external voltage applied and the signal is affected from secondary electron emission, which is known to strongly affect the detection properties in the UV and EUV spectral regions. A different geometry reported is a polycrystalline sandwiched photodiode structure (V.I. Polyakov et al., 1998, L. Thaiyotin et al., 2002) with a contact on the diamond growth surface and a backside contact on the silicon substrate. However, the CVD diamond performance is limited in this case by the polycrystalline structure due to defect states in the band gap introduced by the grain boundaries (R. D. McKeag & R. B. Jackman, 1998, L. Barberini, 2001), which affects the photoelectric properties and alters the detection characteristics. On the other hand, detector grade natural diamonds are extremely rare and expensive, while high pressure high temperature (HPHT) diamonds have their performance strongly worsened by defects and impurities (E. Pace et al., 2000). A great effort is therefore being devoted to produce device-grade Single Crystal Diamond films (SCD) by homoepitaxial CVD growth on low-cost diamond substrates (S. Almaviva et al., 2009, 2010a). A few years ago, at the University of Rome “Tor Vergata” laboratories, CVD single crystal diamond films were used to obtain a new class of detectors with a layered structure. Thanks to the combination of boron doped and intrinsic single crystal diamond films, together with the possibility to easily build Schottky junctions on intrinsic diamond by thermal evaporation of the metal contacts, it has been possible, by using simple multilayered a p-type/nominally intrinsic diamond/metal layered structures, to obtain high quality and highly reproducible devices which can be effectively used for detection (UV and X-rays) photons.
In this chapter, the fabrication and characterization of two Schottky photodiodes, based on p-type/intrinsic diamond/metal (PIM) Schottky junction in two different operative configurations is reported (S. Almaviva et al., 2010b). One detector has been designed in transverse configuration with a semitransparent metallic contact evaporated on the diamond surface, while the second one operates in a planar configuration with interdigitated fingers on the diamond surface. One set of fingers is made of aluminium and the second one is made of p-type diamond.

2. CVD single crystal diamond films production (detector grade)

The reactor for Microwave Plasma Enhanced Chemical Vapour Deposition (MPECVD) diamond growth, used at the Mechanical Engineering Department of the University of Rome “Tor Vergata” laboratories is illustrated in Fig.1. This apparatus is used for the deposition of diamond films in the present work. This deposition chamber was built by developing the necessary parameters to obtain films of diamond having the best physical characteristics, together with a high degree of reproducibility results.

![Fig. 1. General scheme of the apparatus for CVD diamond films deposition](image)

The apparatus for CVD diamond deposition is composed by a tubular microwave CVD reactor and the growth chamber consists essentially in a quartz tube into which the precursory gases flow, controlled by four flowmeters, (a mixture 100 H\textsubscript{2}/1 CH\textsubscript{4} sccm for the intrinsic growth and 90+70 H\textsubscript{2}/ 2 CH\textsubscript{4}/ 10+30 H\textsubscript{2}-B\textsubscript{2}H\textsubscript{6} sccm for the doped one) at a stationary fixed pressure (typically 120+130 mBar).

The tube is put across a waveguide connected to a commercial 2.45 GHz MW generator (Sairem® magnetron, 2kW maximum MW power) and the typical microwave power for both SCD growths is 500+550 W. The quartz tube acts then as a resonant cavity for the microwaves and a stable plasma burns at the centre of the tube, coupling to the sample holder, that is properly connected to a water cooling system, and thus to the diamond substrate on which the homoepitaxial deposition process takes place. The position of the
plasma can be modified through a “plunger” (a sliding short circuit) that terminates the waveguide and its coupling with the substrate can be optimized by means of a 3-stub tuner put between the MW generator and the quartz tube. The temperature of the growing film is monitored in real-time through an infrared optical pyrometer and can be maintained constant during the whole CVD process thanks to a proper software which follows the growth and varies properly the MW power supplied by the generator. In this type of reactor, microwaves give energy to the electrons of the plasma, which, at his time, give energy to the gas through molecular collisions, heating it. The chemical and physical reactions, which follow this heating, allow the formation of carbon that is deposited on diamond substrate. The chemistry of the diamond deposition is described in the next paragraph. It’s worth underline that diamond growth is obtained only if growth parameters (gas composition, pressure, plasma temperature, microwave power density, etc) are chosen in the appropriate way. In particular, the grow rate of diamond with this method vary from 0.2 \( \mu \text{m/h} \) to 10 \( \mu \text{m/h} \), depending on growth parameters and chamber geometry.

3. P-type/ intrinsic diamond/ metal (PIM) Schottky photodiode

Schottky photodiodes are based on synthetic SCD films produced in our laboratories by using MPECVD technique previously explained. The nominally intrinsic diamond is deposited by using a completely separated apparatus in order to avoid any boron contamination. In fact, the presence of impurities in the intrinsic SCD active layer would determine a drastic worsening of the resulting performances, in terms of increasing of the dark current, temporal instability, memory effects, priming, slow response times, worsening of the spectroscopic performances, etc. All CVD films were principally characterized by X-ray diffraction, cathodoluminescence and Scanning Electron Microscopy, confirming the single crystal homoepitaxial deposition and the good crystal quality of the grown samples.

The single crystal p-type diamond was grown keeping constant the microwave power, while the intrinsic layer was grown keeping constant the temperature. The typical growth conditions are reported in Table 1.

<table>
<thead>
<tr>
<th>Typical growth parameters</th>
<th>Intrinsic diamond</th>
<th>p-type diamond</th>
</tr>
</thead>
<tbody>
<tr>
<td>Substrate</td>
<td>(100) HPHT type Ib</td>
<td></td>
</tr>
<tr>
<td>Plasma composition</td>
<td>( \text{CH}_4-\text{H}_2 )</td>
<td>( \text{CH}_4-\text{H}_2-\text{B}_2\text{H}_6 )</td>
</tr>
<tr>
<td>Gas flow rate</td>
<td>1-100 sccm</td>
<td>2-100-10×30 sccm</td>
</tr>
<tr>
<td>Microwave power</td>
<td>500-6000W</td>
<td>~500W</td>
</tr>
<tr>
<td>Temperature</td>
<td>720 °C</td>
<td>450 °C ( \rightarrow ) 650°C</td>
</tr>
<tr>
<td>Pressure</td>
<td>120 mbar</td>
<td>150 mbar</td>
</tr>
<tr>
<td>Thickness rate</td>
<td>1( \mu \text{m/h} )</td>
<td>2( \mu \text{m/h} )</td>
</tr>
</tbody>
</table>

Table 1. Typical growth parameters

The purity of CH4 and H2 gases were 99.9995% and 99.9999% respectively. Boron doping was performed by adding dyborane-hydrogen gas mix (100 ppm \( \text{B}_2\text{H}_6 \) in hydrogen) to the source gases. In the following section, the fabrication process of both photodiodes with different structures is reported.
3.1 Transverse configuration
The diamond photodiode in transverse configuration consists of a multilayered structure obtained by a two-step deposition process. A conductive boron-doped diamond homoepitaxial layer (see Fig. 2 (b)), used as a backing contact, is deposited, at first, by Microwave Plasma Enhanced Chemical Vapour Deposition (MWPECVD) on a commercial low-cost synthetic High Temperature High Pressure (HPHT) <100> type Ib single crystal diamond (SCD) substrate, 4 × 4 × 0.5 mm³ in size and approximately 400 µm thick. The boron concentration was estimated by fitting the Resistivity - Temperature curves, obtaining values in the range 10²⁰ cm⁻³. After that, a nominally intrinsic diamond layer, which operates as the detecting region, is homoepitaxially grown on the doped one (see Fig. 2 (c)). Its thickness can vary from a few microns up to more than 200 microns. Due to the small penetration depth of the UV radiation in the 10–200 nm spectral range (D. Palik., 1991), the detecting region of diamond has a thickness of approximately 2 µm. In the case of the soft X-ray detection, a higher thickness can be chosen (about 30 µm). The nominally intrinsic diamond is deposited by using a completely separated apparatus in order to avoid any boron contamination.

After the growth, each diamond layer has been oxidized by isothermal annealing at 500 °C, for 1h in air, in order to remove the H₂ surface conductive layer. Finally, a circular metal electrode, about 3 mm diameter is deposited on the diamond surface by thermal evaporation and annealed silver paint was utilized in order to provide an almost ohmic contact to the B-doped layer (see Fig. 2 (d)).

Fig. 2. Diamond deposition process of the PIM detector
Such a structure acts as a p-type/ intrinsic/ metal (PIM) Schottky Barrier Diode: the metal creates a Schottky junction with the intrinsic diamond, which acts as a drift-layer, and the SCD p-type layer, holes injector, determines the unipolarity of the device under direct polarization. The device, therefore, operates in typically reverse biased mode, with a negative voltage on the boron doped contact and the metal top contact grounded.

3.2 Planar configuration
In this section, it is report on the fabrication process of extreme ultraviolet (EUV) photovoltaic single crystal diamond Schottky diodes developed at the University of Rome “Tor Vergata” operates in planar configuration with an interdigitated contact structure on the growth surface of the intrinsic
diamond layer. One set of fingers is made of aluminium and the second one is made of p+-type diamond. Two steps of a standard photolithographic technique are used for the fabrication process (see Fig.3) for the diamond detector with interdigitated finger electrodes (IDT-PIM in the following). First, an intrinsic diamond layer is homoepitaxially grown by MWPECVD on a commercial HPHT single crystal diamond substrate. As previously mentioned, annealing in air is employed in order to remove the surface conductive layer of the as-grown diamond film.

After the annealing process, p-type diamond interdigitated fingers are selectively grown on the top of the intrinsic diamond layer by using a patterned Cr plasma-resistant coplanar mask (Fig.3 (a)-(d)). After removal of the chromium mask by wet etching, the interdigitated Al electrode is fabricated by using a second mask which is aligned to the pattern previously obtained.

The Al fingers are patterned by standard lift-off photo-lithographic technique and by thermal evaporation on the CVD intrinsic diamond surface (Fig.3 (e)-(g)). The width and the gap between two fingers are both 20µm. A SEM image of the IDT-PIM device in Fig.3. The IDT-PIM is simply mounted in an sample holder for UV measurements. In this case, the measured photocurrent of IDT-PIM detector can contain both photoemission current and photoconductive current. The photoemission contribution contains electron emission arising from Al fingers and from p-type diamond exposed to the UV irradiation.

![Fabrication process of the IDT-PIM photodetector and SEM image of the IDT-PIM device: (1) p-type diamond, (2) intrinsic diamond and (3) Al contact.](image)

**4. Electrical characterization**

The investigation of the physical principle of operation of such photodiode through the study of their electrical detection properties is reported in this paragraph. The physics of the devices is clearly based on the existence of a Schottky barrier, as demonstrated by the fact that the detectors are able to operate even with no external bias applied. The aim of the electrical characterization of the devices is, therefore, to analyze the Schottky barrier parameters, measured using the I-V curves from which the barrier height can be extracted. Moreover, the study of metal/p-type diamond contacts to obtain a good ohmic contact is also reported.
4.1 Ohmic contact on p-type diamond

High quality ohmic contacts are one of the most frequently encountered problems in the development of the right material for electronic devices applications. The properties of such electrical contacts directly contribute to the active devices performance. In general the requirements for ohmic contacts can be summarized as follows (M. Werner, 2003):

- low contact resistivity
- good adhesion
- high thermal stability
- high corrosion resistance
- bondable top-layer
- about zero voltage drop

As the low doped diamond films exhibited semiconducting behaviour, it was necessary to study the metal/diamond contacts to verify if they obey Ohm’s law. Most contacts to common semiconductors are depletion contacts which result mainly from the action of surface states. They can show, however, an ohmic behaviour with a linear current–voltage characteristic on degenerated doped semiconductors. In the case of a depletion contact, the contact resistivity varies exponentially with the Schottky barrier height. The ohmic behaviour of the depletion contact can be achieved either when the barrier height is small, so that the charge carriers can easily overcome the barrier (thermionic emission) or when the charge carriers are able to surmount the depletion region by quantum-mechanical tunnelling.

![Graph](image)

Fig. 4. I-V characteristic of annealed silver paint/p⁺-type diamond junction

The depletion layer width of a metal/semiconductor contact is proportional to the square root of the reciprocal doping concentration \( W \sim N_A^{-1/2} \) (S.M. Sze, 1981). Consequently, the
depletion layer width decreases with the increase in doping concentration and the tunnelling probability increases. So a good ohmic contacts are obtained by heavily doping the p-type diamond layer (doping levels much larger than $10^{20}\text{cm}^{-3}$). The resulting layer $p^*$, which is highly doped by B, was metalized by silver paint annealed at 500°C for 10 min. The I-V characteristic is reported in Fig.4 where is also reported the specific resistance calculated by ohm’s law.

4.2 Shottky contact on intrinsic diamond layer

The electrical characterization of the metal/intrinsic diamond Schottky junction of the devices was performed at room temperature in a vacuum chamber with a background pressure of $10^{-4}$ mbar by measuring the current-voltage (I-V) characteristics by using a Keithley 6517A pico-amperometer.

The I-V characteristic was obtained by applying a voltage to the metal contact while the p-type diamond layer is earthing. Fig.5 shows the typical I-V characteristic of the diamond Schottky photodiodes. When the p-type rectifying contact is reverse biased by connecting the metal to positive terminal, holes are repelled from the interface and the bands are away bent down. The potential barrier for the holes is increase, as is the width of the depletion region. The resulting net current is very low (reverse biased). If instead the metal is connected to the negative terminal, then forward biasing results as the holes are attracted toward the metal interface (forward biased).

![Fig. 5. Typical I-V characteristic of the PIM device](image-url)

In Fig.5, it’s clearly seen the different behaviour of reverse and forward current. When a negative voltage (forward voltage) is applied on the metal electrode a hole current starts flowing from the p-type diamond, via the nominally intrinsic diamond region, towards the Schottky contact. The rectification behaviour of the both photodiodes is observed with a very high rectification ratio of about $10^8$ at ±3V. For values of $|V_B| < |V_{on}|$, where $V_{on}$ is “turn- on voltage” that in figure is about - 1 V , the forward current is due to generation-recombination effects and leakage superficial current and it’s similar at the reverse current. Increasing the forward bias, in the region between approximately -1 V and -1.6 V forward voltage ($V_F$), the current rises exponentially with $V_F$.

In this region the forward current density ($J_F$) is well described by the thermionic emission (TE) theory. The thermionic emission theory by Bethe is derived from the assumptions that...
the barrier height is much larger than kT, thermal equilibrium is established at the plane that determines emission, and the existence of a net current flow does not affect this equilibrium. Bethe’s criterion for the slope of the barrier is that the barrier must decrease by more than kT over a distance equal to the scattering length. The resulting current flow will depend only on the barrier height and not on the width, and the saturation current is not dependent on the applied bias. Then the current density of majority carriers from the semiconductor over the potential barrier into the metal is expressed as (M. Brezeanu et al., 2007):

\[ J_F = J_S \left[ \exp \left( \frac{qV_F}{nkT} \right) - 1 \right] \]
\[ J_S = A^*T^2 \exp \left( \frac{-q\Phi_{\text{BI}}}{kT} \right) \]  
(1)
\[ A^* = \frac{m_p^*}{m_0} A^0 \]

where \( n \) is the ideality factor (\( n \geq 1 \) and it informs the experimental I-V characteristic deviates from the behaviour SBD ideal (\( n = 1 \)), T the absolute temperature (Kelvin), k the Boltzmann’s constant, \( J_S \) the saturation current density, \( A_0 \) the Richardson’s constant (120.173Acm\(^{-2}\)K\(^{-2}\)), \( A^* \) the Richardson’s effective constant, \( m_0 \) and \( m_p^* \) electron mass and effective mass hole in diamond (\( m_p^* = 0.7 m_0 \)) and \( \Phi_{\text{BI}} \) the Schottky barrier height. From the exponential fit of the I-V characteristic, it is possible to estimate the saturation current density \( J_S \) and the ideality factor \( n \). Substituting the values obtained from the fit in the following equation

\[ \Phi_{\text{BI}} = \frac{kqT}{q} \ln \left( \frac{A^*T^2}{J_S} \right) \]  
(2)

it’s possible estimate the Schottky barrier height. The values obtained for IDT-PIM and PIM photodiodes are 1.65 eV and 1.8 eV respectively.

5. Extreme UV characterization

The photodiodes have been tested over the extreme UV spectral region from 20 to 120 nm, using He and He Ne DC gas discharge as radiation sources and a toroidal grating vacuum monochromator (Jobin Yvon model LHT 30) with a 5Å wavelength resolution. The dimension of the optical aperture is 0.25 × 6.00 mm\(^2\); a manual shutter is used to switch on and off the UV radiation. The experimental apparatus of UV characterization is reported in the following picture.

The photoresponse measurements have been performed in a vacuum chamber, at a pressure of 0.03 mbar. By using a three (X-Y-Z) dimension mechanical stage powered by stepper motors, it is possible to locate the photodetector in front of the beam light and to compare its response with that of a calibrated NIST silicon photodiode (http://www.ird-inc.com) placed in the same position, which measures the absolute photon flux. A raster scansion of the beam light was performed on the detector surface so to position the photodetectors where their response has a maximum (see Fig.6(b)).
A hole, 2 mm in diameter, is used to collimate the radiation on the sensitive area of the detectors and to obtain the same illuminated area on the silicon photodiode. The photocurrent is measured by an electrometer (Keithley 6517A), using the internal voltage source. Because of different geometry adopted by the two devices, they are measured differently. The PIM detector is encapsulated in a copper/vetronite shielded housing with a 2 mm pinhole. In such housing, the Al contact is grounded and the photocurrent is measured from p-type diamond so that the signal is not affected by the eventual presence of secondary electron emission current from the illuminated contact.
The IDT-PIM is simply mounted in an sample holder for UV measurements with the same 2 mm pinhole. In this case, the measured photocurrent of IDT-PIM detector can contain both photoemission current and photoconductive current. The photoemission contribution contains electron emission arising from Al fingers and from p-type diamond exposed to the UV irradiation. The typical current – voltage (I-V) characteristics in dark current and under irradiation have been measured at room temperature of two detectors are shown in Fig. 7. The devices operate in the reverse bias mode because when operating in the forward bias mode, the photocurrent is masked by the dark current. The dark current is very low (<0.1 pA) below about +10 V, as expected for a metal/diamond rectifying contact. The photocurrent vs. applied voltage is also reported in the same figure when the device is exposed to UV radiation and 30.4 nm (He lines) and 73 nm (Ne line). The device shows a photocurrent response even at zero voltage bias, exploiting the internal junction electric field. The photocurrent is almost constant with increasing positive voltage, while the dark current increases by about two orders of magnitude. Remarkably, thus, the best signal-to-dark current (SDR) ratio (see Fig. 7 (b)) is obtained at zero bias voltage, so that in the following, the devices have been operated with no external bias voltage applied.

5.1 Temporal response
Temporal response measurements upon exposure to UV radiation have been performed according to the following procedure: at first the dark current value was recorded for several seconds keeping the light shutter closed, until the steady state value had been reached; then the shutter was opened and the photocurrent was measured. Finally, the shutter was closed again until the dark current reached the initial value, before starting a new measurement run. The detectors time response, upon exposure to UV radiation, have been measured by opening and closing a manual shutter during the acquisition. The temporal response of the tested devices is reported in Fig. 8 (a) under UV illumination of the He-Ne DC gas discharge radiation source.

![Fig. 8. a) Temporal responses under illumination of He-Ne DC gas discharge radiation source. b) The magnification of fall time of the both devices.](www.intechopen.com)
The response is reproducible and no undesired effects such as persistent photocurrent and priming or memory effects, which are often observed in diamond UV detectors (C. E. Nebel et al., 2000, A. De Sio et al., 2005, M. Liao et al., 2008), are observed. negligible. However, it is obtained only after the very first irradiation: the device, just mounted, reaches the described performance only after a pre-irradiation time of few minutes Fig.8 (b) shows rise and fall times of the signal of about 60 ms, which corresponds to the acquisition rate of the used electronic chain.

5.2 Linearity

A useful detector is expected to exhibit linear response with photon flux, i.e. a constant responsivity up to a saturation point where space charge effects prevail and no more electron-hole pairs can be collected under illumination. The calibration of linear detectors and related electronics is much simpler. The linearity of the photodetectors have been investigated varying the current intensity of the plasma. The photocurrent \( I_{ph} \) measured vs. the incident optical power \( P \), under irradiation of He-Ne gas discharge radiation is shown in Fig.9. We used a power law: \( I_{ph} = A + B \cdot P^C \) to fit the data. Here \( A \) is the offset corresponding to the dark current, \( B \) is the photosensitivity (provided \( C=1 \)) and \( C \) is a linearity coefficient. The graph shows the measured data as well as the fitting function. In this spectral region, both photodetectors shows remarkably good linearity, \( C \) being 1 in all cases within the error.

![Fig. 9. Linearity of IDT-PIM and PIM photodiodes.](image)

5.3 Extreme UV spectroscopy

The normalized emission spectra of a DC discharge He and He-Ne lamp measured in unbiased mode by the detectors are reported in Fig.10. All spectral lines are clearly resolved and observed with a good signal to noise ratio, demonstrating the high photodetection capabilities of the CVD single crystal diamond grown in the extreme UV spectral region. All spectral lines are classified by the NIST Atomic Spectra Database Lines Form from the following website: http://physics.nist.gov/PhysRefData/ASD/lines_form.html.
Fig. 10. He-Ne emission spectrum measured by the two devices.

In particular, the low intensity lines of the He-Ne spectrum in the wavelength range 20-30 nm are easily resolved by PIM detector.

Fig. 11. He-Ne spectrum measured by PIM detector in the range 20-30 nm.

5.4 Responsivity and external quantum efficiency

The absolute spectral response of the PIM detectors is measured by comparison with a calibrated photodiode exposed to the same source on the same optical area of about 1 mm². The spectral responsivity, expressed in amperes per watt (A/W), is defined as the photocurrent per unit incident optical power and can be evaluated from the relationship

\[
R_d = \frac{R_{Si} I_d}{I_{Si}}
\]

where \( R_{Si} \) is the responsivity of the calibrated silicon photodiode at a given wavelength, \( I_{Si} \) and \( I_d \) are the photocurrents measured by the silicon photodiode and the diamond detector, respectively.

The responsivities of both photodiodes are reported in Fig.12. The responsivity of the PIM device decreases monotonically as the wavelength increases until about 80 nm while at 120 nm...
nm an increased value is observed. At 98 nm the signal is below the noise level so that only an upper limit can be provided. However, the presence of a minimum in the responsivity around 100 nm can be clearly deduced from Fig.12.

![Responsivity of the both devices](image)

**Fig. 12. Responsivity of the both devices.**

The responsivity of the IDT-PIM detector is much lower than that of the PIM detector at short wavelength (below 50 nm) showing a maximum at about 73 nm. The increased sensitivity of the IDT-PIM device at intermediate wavelength could be probably ascribed to the contribution of photoemission current as already reported in the literature (T. Saito et al, 2006). For both the devices the absolute responsivity measured at around 50 nm is comparable to the best results reported in the literature for diamond based EUV detectors (A. BenMoussa et al., 2006).

The External Quantum Efficiency (EQE) spectrum, estimated by: \[ EQE = \frac{1240 \cdot R_d}{\lambda} \] is reported in Fig.7 for the PIM devices.

As mentioned above, the photocurrent measured by IDT-PIM detector includes the contains both photoconductive current and photoemission current, arising from secondary electron escape from Al fingers, which also depends on the wavelength (J. Ristein et al, 2005, W. Pong et al., 1970). On the contrary, in the encapsulated PIM device the illuminated contact is grounded and the current flowing from the boron doped layer is not affected by secondary electrons contribution. Moreover, the more homogeneous electric field configuration of the PIM device allows a simple analysis of the detection process.

In order to investigate the effect of the metallic Schottky contact upon the detection performance of the PIM devices, different semitransparent metals (thickness < 10nm) have been thermally evaporated on the oxidized surface of single crystal CVD intrinsic diamond layers.
Fig. 13. External quantum efficiency EQE of the two photodiodes between 20 and 120 nm.

The absolute spectra responsivity curves versus different metal contacts of the devices are shown in Fig. 14. All the devices have a maximum of the responsivity at lower wavelengths and a sharp cutting edge for longer wavelengths while at around 120 nm an increased value is observed. The lowest responsivity, between 50 - 100 nm, has been measured for the device having Cr as an electrode. The device having Ag and Pt contacts shows rather similar trend of the responsivity, whereas Al contact shows the best results in the UV performances.

Fig. 14. External quantum efficiency EQE of the PIM devices between 20 and 120 nm as a function of the type of the metallic contact.
5.5 UV/visible rejection ratio
The photoconductive response was tested over a wide spectral range, extending from the extreme UV (EUV) up to the visible. The 210–500 nm range was investigated using an Optical Parametric Oscillator (OPO) 5 ns pulsed laser (Opolette laser by Opotek). The laser beam was scattered by an optical diffuser in order to prevent signal saturation of the electronic chain and the diamond detector was placed 10 cm away from the diffuser. A 500 MHz Le Croy WaveRunner 6050 digital oscilloscope was used to acquire the output signal.

Two different connection configurations were used:
1. Direct recording of the detector output by the digital oscilloscope
2. Integrated measurement by an Ortec142A charge preamplifier.

The signal provided by a pyroelectric power meter was used to normalize the diamond detector output, in order to take into account the wavelength dependence of laser pulse amplitude and the intrinsic fluctuations of the beam intensity.

The visible-blind properties of the photodetectors were tested by measuring the photoresponse at different wavelengths in the 210–500 nm range.

In Fig.16 (a) the device responsivity of the PIM detector is reported as a function of the incident laser radiation wavelength, normalized to the pyroelectric power meter signal. A 3 orders of magnitude variation was measured when moving across the band gap wavelength of 225 nm. Such a drop increases up to 5 orders of magnitude when the UV to visible rejection ratio is considered. It should be stressed that a very stable and reproducible response was observed in the whole energy range and irradiation memory or pumping effects were not observed.

In addition, a linear increase in the photoresponse as a function of calculated radiation intensity was observed measuring the output signal at decreasing device distances from the optical diffuser.

The time response at 220 nm of the investigated PIM detector is reported in Fig. 16 (b). As clearly seen in the Fig.16 (b), the device response to a laser pulse at 220 nm, measured through a bias Tee and recording by the digital oscilloscope (Le Croy 500MHz), shows an exponential decay time constant of about 100ns. The reason of this trend of output response is due to electrical circuit of the device. In fact, an RC circuit, the value of the time constant is equal to the product of the circuit resistance and the circuit capacitance. Therefore, taking into account the depletion capacitance measured by C-V curves of about 100pF and the resistance of p-type diamond film ~1kΩ, the time constant result to be $\tau = 100\text{ns}$. 

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Fig. 16. a) Normalized responsivity of PIM device as a function of the incident laser radiation wavelength. b) The device response to laser pulses directly obtained by the digital oscilloscope.

The visible-blind properties of the IDT-PIM device were also tested by measuring the photoresponse at different wavelengths in the 210–500 nm range. In this region, the spectral response shows a visible/UV rejection ratio of about 4/5 orders of magnitude, as clearly seen in Fig. 17(a). Moreover, the time response at 220 nm of the investigated detector is reported in Fig. 17(b). The Fig. 17(b) shows the device response to a laser pulse at 220 nm, which have a full width at half maximum (FWHM) of about 25 ns, and the time response is faster than that of PIM detector. In fact, in this case, the parallel capacitance of the photodiode is very low, about 15pF. Interdigitated structure, therefore, can be optimized in order to build a ultrafast XUV detector, for time resolution.

Fig. 17. a) Normalized responsivity of IDT-PIM device as a function of the incident laser radiation wavelength. b) The device response to laser pulses directly obtained by the digital oscilloscope.

6. Conclusion

Two detectors were fabricated at the University of Rome “Tor Vergata” with a structure that acts as a metal/intrinsic/p-doped diamond photovoltaic Schottky diode. The two detectors
operate in different configurations: one in transverse geometry and the other one in planar configuration.

We have measured the electrical characteristics and tested the performance under continuous vacuum UV photon irradiation of the two devices. A general result of our experiments is that diamond detectors are very sensitive devices showing a very low dark current and very good signal-to-noise ratio. The responses are reproducible and undesired effects such as persistent photocurrent, priming or memory effects are negligible for both devices. The response time could be very fast and it is much lower than the acquisition rate of the used electronic chain (≈ 60 ms). These results indicate the high quality of our CVD diamond grown for UV applications.

The responsivity and the EQE of the two devices show an opposite behaviour as a function of the radiation wavelengths due to the different operative configurations. In particular the PIM detector is more efficient at lower wavelengths and present a drop of sensitivity at approximately 100 nm. The IDT-PIM is less efficient at low wavelength and has a maximum efficiency at about 74 nm.

The visible-blind properties of the photodetector were also tested by measuring the photoresponse at different wavelengths in the 210–500 nm range. A 3/4 orders of magnitude variation was measured by diamond based detectors when moving across the band gap wavelength of 225 nm. Moreover, the spectral response shows a visible/UV rejection ratio of about 5 orders of magnitude for both photodiodes. Finally, the device response to laser pulses at 220 nm is different in two cases due to the different electrical circuit of the two devices. In particular, the time response of IDT-PIM detector is faster than that of PIM detector.

7. Acknowledgment

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8. References


Photodiodes or photodetectors are in one boat with our human race. Efforts of people in related fields are contained in this book. This book would be valuable to those who want to obtain knowledge and inspiration in the related area.

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