Review



Review on advances in microcrystalline, nanocrystalline and ultrananocrystalline diamond films-based micro/nano-electromechanical systems technologies

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Received: 24 October 2020 Accepted: 14 December 2020 Published online: 25 January 2021

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ABSTRACT

A comprehensive review is presented on the advances achieved in past years on fundamental and applied materials science of diamond films and engineering to integrate them into new generations of microelectromechanical system (MEMS) and nanoelectromechanical systems (NEMS). Specifically, the review focuses on describing the fundamental science performed to develop thin film synthesis processes and the characterization of chemical, mechanical, tribological and electronic properties of microcrystalline diamond, nanocrystalline diamond and ultrananocrystalline diamond films technologies, and the research and development focused on the integration of the diamond films with other film-based materials. The review includes both theoretical and experimental work focused on optimizing the films synthesis and the resulting properties to achieve the best possible MEMS/NEMS devices performance to produce new generation of MEMS/NEMS external environmental sensors and energy generation devices, human body implantable biosensors and energy generation devices, electron field emission devices and many more MEMS/NEMS devices, to produce transformational positive impact on the way and quality of life of people worldwide.

Introduction

Relatively medium level appropriate mechanical and excellent electrical properties, the latter optimized for many years of development of Si microelectronics, prompted researchers to perform R&D to investigate the applicability of Si to fabricate microelectromechanical system (MEMS) devices. The idea of using Si for production of MEMS devices arose from the hypothesis that the MEMS structures could be integrated with the well-developed Si-based electronics

Handling Editor: David Cann.

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circuit/devices on the same Si substrate, providing the basis for fabrication of compact/efficient/low energy consumption Si-based MEMS devices. However, Si has material limitations from the mechanical and surface tribological point of view [1], which are critical properties for MEMS devices. Specifically, Si exhibits both relatively low Young's modulus and fracture toughness, compared to many other materials, and also exhibits comparatively large coefficient of friction (COF) and mechanical wear, resulting in Si-based MEMS devices with high energy loss, due high friction of MEMS devices sliding parts. In addition, Si-based MEMS devices involving components subject to rolling or sliding motion exhibit failure due to wear of components in short time [2–4]. The problems described above can be enhanced by interacting surfaces of components in humid environments, which result in surfaces adhesion to each other [5] or by low device operating temperature and/or heat dissipation induced by a small energy band gap [3]. Therefore, the information presented above indicate that Si is not an appropriate material for fabrication of several MEMS devices, even if they operate at room temperature. On the other hand, research by independent groups revealed that Si MEMS components coated with Si₃N₄ or SiC films exhibit improved mechanical behavior than pure Si in MEMS devices operating at room temperature, However, research by independent groups demonstrated that Si₃N₄ or SiC-coated Si MEMS components also exhibit materials and device fabrication limitations [6, 7].

The information presented above induced scientist and engineers to investigate new materials for fabrication of MEMS devices. In this sense, diamond and diamond-like carbon (DLC) are considered as materials with order of magnitude superior mechanical and tribological properties than SI for MEMS applications. In addition, metals, other than Si semiconductors, polymers, oxide materials, the latter with piezoelectric properties, are being investigated for their suitability for integration with the diamond or DLC materials to provide mechanical action properties. Recently, the world first RF-MEMS switch made of single crystal diamond (SCD) was reported, showing better functionality, including device reliability, lifetime, speed, and electrical performance, compared to existing RF-MEMS switches [8]. Although SCD demonstrated the superiority of this material for MEMS devices, a problem is that SCD is

currently not suitable for industrial/low cost process on large scale substrates, as currently used in the fabrication of Si electronics devises on 300 mm diameter Si substrates. Therefore, polycrystalline diamond (PCD) and DLC films are investigated for fabrication of MEMS devices because they not only exhibit many of the properties of SCD, but they can be grown on a wide range of large area substrates with good film growth rates and thickness and structure uniformity, making them suitable for very large-scale production to fabricate diamond filmbased MEMS devise on large area substrates at low cost. Three distinct forms of PCD are microcrystalline diamond (MCD- grain sizes in the range ≥ 1 micron), nanocrystalline diamond (NCD-grain sizes in the range 10-100s nm) and ultrananocrystalline diamond (UNCD-grain sizes in the range 3-5 nm)). All films mentioned above are grown inside chambers where air has been evacuated, to produce vacuums in the range 10^{-7} – 10^{-8} Torr, and flowing gas mixtures as described in the following sentences. MCD films are grown flowing H₂ (99%)/CH₄ (1%) gases into air evacuated chamber and coupling microwave power to produce a plasma that generate CHx (x = 1, 2, 3) and C species, which upon landing on the substrate surface grow the film [4]. NCD films are grown flowing mixtures of H₂ (96%) /CH₄ (4%) [10]. UNCD films are grown using a unique patented Ar (99%)/ CH_4 (1%) gas mixture that induces the production of C_2 dimers + CHx (x = 1, 2, 3) species, which produce the 3–5 nm grains that define UNCD [2, 4]. DLC films exhibit an amorphous structure displaying some of the typical properties of diamond. DLC exists in seven different forms, all containing significant amounts of sp^3 hybridized carbon atoms bonds (bond of diamond). The reason that there are different types of DLC forms is because even diamond can be found in two crystalline polytypes. DLC has the highest packed atomic structure of all materials involving the diamond phase [9], and it can even be grown at very low temperatures (near room temperature) by pulse laser deposition [10] or ion beam sputter-deposition [11]. MCD to UNCD and DLC films exhibit a number of excellent properties far superior to those of Si material and other materials being used for MEMS and NEMS devices. However, even among the PCD films, there are substantial differences in properties relevant for MEMS and NEMS devices. For example, the COF of MCD is in the range of 0.5–0.3, NCD in the range 0.2–0.3, DLC \sim 0.2, while UNCD exhibits the lowest COF (0.02-0.05) of all PCD materials described above (see review in ref. [4]). All PCD offer high resistance to wear [4]. In addition, two key mechanical properties of diamond and DLC films, particularly superior than for any other materials for MEMS/NEMS devices, containing components with sliding interfaces, are very high Young's modulus and tensile and fracture strength, which make diamond and DLC films particularly suitable for high frequency MEMS/NEMS devices. Diamond and DLC films are also chemically inert against common environmental conditions and are stable in air up to 600 °C, which make them attractive materials for operation at high temperatures above 300 °C, where conventional devices cannot operate. MCD has fairly high thermal conductivity (\sim 1990 W/K m [4]), which is close to the thermal conductivity of SCD ($\sim 2100 \text{ W/K}$ m), the highest of any material both in the direction perpendicular and parallel to the surface of the material [4]. On the contrary, UNCD has a low thermal conductivity, due to the large grain boundary network [4], which makes it an ideal thermal insulation layer. All PCD films exhibit low thermal expansion coefficient, making them suitable for implementation in integrated portable lab on micro-electronic devices, packages or supporting material with high heat dissipation suitable for MEMS thermal actuators with extensive mechanical motion. Diamond and DLC films are the best biocompatible materials because they are made of C atoms (element of life in the human DNA, cells, and molecules), and they are mechanically strong and tribologically efficient, making them the best coatings for implantable medical devices. The surface of diamond and DLC films can be modified to facilitate attachment of specific biological species, providing the bases for a new generation of superior biosensors. Diamond and DLC are also excellent materials for field emission display which is brighter and faster than LCD and will become the ultimate display in the future. DLC is the only material that is fully thermionic, i.e. it can convert heat into electricity, which makes DLC potentially the best solar cell material with projected energy conversion efficiency over 50% [9]. It is worth mentioning that the composite of UNCD and carbon nanotube (CNT) is also a promising material for future thermoelectrical material. Furthermore, diamond also provides a high breakdown voltage for a high-power generation and a low dielectric constant for a better isolation and

coupling characteristics. Based on the information presented above, MEMS/NEMS devices based on diamond or DLC films would provide straightforward integration with diamond electronics in a unified support platform. In addition, diamond filmbased MEMS/NEMS devices will have the added advantage of efficient high thermal controlling properties, relevant for high temperature performing MEMS/NEMS devices.

History of chemical vapor deposition (CVD) of diamond

Natural diamond gems were first inserted as royal jewels in India, while the Romans used them to engrave sapphires. Diamond was identified as a Carbon material in 1796 but was not synthesized out of nature until 1954. In the late 1960s, two separate groups, one from the Soviet Union led by Boris Deryagin (1970) and the other from the United States headed by John Angus (1968), independently showed that diamond films could be grown on surfaces of materials seeded with MCD particles. MCD films were grown by chemical vapor deposition at low pressures from a vapor containing hydrocarbons (CH₄) and hydrocarbon (CH₄)-hydrogen (H) gases mixtures [12, 13]. Angus and co-works were the first to report the critical role of atomic hydrogen (H) in achieving metastable diamond film growth, such that H atoms were identified as a preferential etchant for removing a graphite impurity face against the diamond face. due to H atoms more efficient chemical reaction with sp^2 open bonds of C atoms in graphite. On the other hand, Deryagin and Fedoseev demonstrated the growth of diamond films on non-diamond substrates made of material such as metals and silicon, which provided the pathway for intensive research to develop processes to produce low cost diamond coatings in the 1980s. The key gases used to grow the MCD and NCD films have been CH4 and H₂, generating CHx^{+ and H} (x = 1, 2, 3) and H^{* and H} species via cracking on the surface metal (mainly tungsten (W) filaments heated to $\sim 2200 \,^{\circ}\text{C}$, via electric current, or cracked by microwave power applied to the gas mixture, producing a plasma (mixture of ionized (H⁺) and neutral (H^o) and ionized molecules CH_x^+ (x = 1, 2, 3) and neutrals CHx^o , to induce carbon (C) atoms mobility on the substrates' surface, so C atoms bond to each other with



the sp^3 diamond bond, while simultaneously H⁺ ions and H^o atoms etch the graphitic phase that grows as an impurity phase. The grown films were mainly formed by randomly oriented MCD grains. The problem related to MCD films is that the surface is very rough (\geq 1-micron rms roughness), which is not appropriate for most MEMS/NEMS devices involving parts sliding upon each other.

The next significant step in the development of diamond films grown by the CVD process took place in Japan. A team led by Nobuo Setaka at the National Institute for Research in Inorganic Materials (NIRIM), Tsukuba, developed methods for the rapid growth of diamond films at low pressures. This work started in 1974 and by 1981, the NIRIM group had published a large number of scientific papers documenting their success in growing diamond films at rates of up to 1 μ m per hour [14]. The stunning achievements of the Japanese researchers rekindled commercial interest in CVD diamond films, particularly in the USA, where, by the end of the 1980s more than 30 companies were investigating the possibilities of this new diamond material and how it could be applied to their business.

UNCD films, discovered in the early 1990s at Argonne National Laboratory, exhibit specific superior properties over traditional MCD and NCD films. The history of the discovery and characterization is much less well known and described in detail in [4, 15]. Along with the development of the CVD diamond deposition technique, DLC has also been relatively developed for low temperature coating which offers low friction and wear properties relevant to sliding, rolling or rotating parts in MEMS/ NEMS devices. The first report of DLC films was by Aisenberg and Chabot [16] in 1971, followed by a series of reports by Holland [17] and Weissmantel [18, 19]. Details of wok on DLC films will not be discussed in this paper.

Material properties

Diamond properties

Diamond has emerged as a transformational material to enable a new generation of MEMS?NEMS devices, due to its excellent complementary mechanical, tribological, chemical, physical, and electronic properties. Diamond is a carbon allotrope with the highest density of any carbon-based material featuring a cubic crystal structure involving two interpenetrating face-centered cubic (FCC) lattices, separated from each other by 1/4 of a lattice diagonal. Each C atom is tetrahedrally coordinated, via strong, covalent bonds (called sp^3 bonds) to four neighbor C atoms [4, 13, 14], as shown in Fig. 1. The covalent bonds between carbon atoms in diamond is called covalent sp^3 -bonding. The bond length and lattice constant are 1.54 and 3.56 angstroms, respectively [20].

The outstanding multifunctional properties of diamond stem from the super strong/tight sp^3 bonds. The large activation barrier between the sp^2 and sp^3 bonding states induce the stable diamond structure, although the most thermodynamically stable microstructure for carbon is graphite. In addition, diamond exhibits a superior wide bandgap, making it an outstanding semiconductor for electronic and optical devices, such as ultraviolet light emitting diodes, cold cathode electron emitters, and high-power and high-frequency devices.

Diamond's well-known properties include the highest hardness compared to any other material, the highest thermal conductivity both perpendicular and parallel to the surface of the solid, and optical transparency, in pure form. In its natural or undoped form, diamond is an excellent electrical insulator with a bandgap of 5.45 eV, as shown in fig. 2 [21–23].

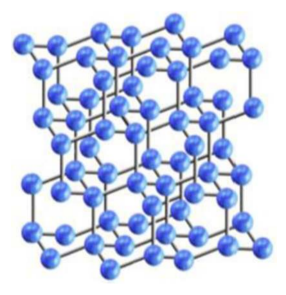


Figure 1 Diamond crystalline unit cell with the electronic structure (four sp^3 orbitals) of one of the atoms marked [9].

Conduction band minima occur along <100> direction at nearly 0.7 of the distance to the edge of the Brillouin's zone. The valence band maximum occurs at the center of the Brillouin's zone, Γ . Thus, the minimum bandgap in diamond is indirect. When doped with boron (B) atoms, diamond exhibits semiconducting or even semi-metallic electrical properties. The properties of natural diamond, compared with other semiconductor materials explored for MEMS/NEMS devices, are summarized in Table 1

Different forms of diamond

There are several types of diamonds for different applications, including natural diamond, synthetic industrial diamond, SCD and polycrystalline diamond (PCD) and diamond-like carbon (DLC) films, with films produced by chemical vapor deposition (CVD) and physical vapor deposition (PVD) methods. Natural diamond has been classified historically by the nature and concentration of impurities. Almost all diamond, natural or synthetic, always contains impurities such as N or B atoms. Natural diamonds are classified by types Ia, Ib, IIa and IIb, based on the quantity of impurities found within them, as shown in Table 2 [24]. Ia is the most common type of natural diamond, with up to 0.3% of N atoms. All other three types are very few in nature.

Artificial diamonds are produced by a High Pressure/High Temperature (HPHT) process, whereby graphite and a metallic catalyst are inserted in a hydraulic press and subjected to a combined high temperature/pressures process. Graphite converts into diamond in a few hours. Most industrial diamonds are type Ib, containing up to 5 ppm of N atoms. Both natural diamond and synthetic diamond are SCD. Considering the high cost, HPHT synthetic condition and difficulty of processing, these two types of diamonds are of very limited use for fabrication of MEMS devices. Instead, PCD films, produced mainly by the CVD process, can provide remarkable properties for high-performance MEMS/ NEMS devices, with relatively low fabrication cost.

Three types of PCD thin films, featuring different microstructure, surface morphology, and properties, have been produced, and their properties characterized in systematic studies. Standard processes to grow PCD films, include microwave plasma chemical vapor deposition (MPCVD), hot filament chemical vapor deposition (HFCVD) or any other CVD methods, The processes described above involve flowing hydrogen-rich gas mixtures $(CH_4/H_2 = 0.1-4\%)$ [25-28, 29-32], into the MPCVD, HFCVD, or other CVD systems, and coupling the power to produce plasmas, which result in the growth of microcrystalline diamond (MCD) (typically 1-10 µm grains) with columnar structures as shown in Fig. 3a, and nanocrystalline diamond (NCD) (typically 50–100 nm grains), as shown in Fig. 3b [318]. NCD films are defined by their grain sizes (10–100 s of nm) and relatively high nucleation density, as demonstrated by Philip et al. [28], Sekaric et al. [29], and Fuentes et al. [318], and they do not exhibit cauliflower type structures characteristics of films grown with high CH₄ levels or with reduce atomic hydrogen

Properties	Si	GaAs	SiC	Natural Diamond
Density (g/cm ³)	2.329	5.317	3.216	3.52
Melting point (°C)	1412	1240	2540	3827
Hardness (GPa)	8.5	7	24.8	100
Young's modulus (GPa)	130-180	85	700	1050-1200
Poisson's ratio	0.22-0.24	0.31-0.32		0.1-0.21
Lattice constant (Å)	5.43	5.65	4.36	3.57
Band Gap (eV)	1.12	1.42	3.0	5.45
Carrier mobility Electron (cm ² /V·s)	1450	8500	400	1800-2000
Hole $(cm^2/V \cdot s)$	500	400	50	1600-2100
Dielectric constant	11.7	10.9	10	5.7
Breakdown voltage (× 10^6 V/cm)	0.37-0.5	0.6	2–3	4–20
Intrinsic resistivity (Ω ·cm)	1×10^{3}	1×10^{8}		1×10^{16}
Thermal conductivity (W/cm·K)	1.5	0.5	5	20
Thermal expansion coef. (× $10^{-6}/{^{\circ}C}$)	2.6	5.9	4.7	1.1

Table 1Comparison ofdiamond properties with othersemiconductors [23]



Table 2 Diamond

classification according to the amount of lattice impurity [24]

Diamond Type	Nitrogen (ppm)	Boron (ppm)	Color
Ia	2000	_	Clear to yellow
Ib	100-1000	_	Green, brown, yellow
Ib	1-100	_	Yellow
IIa	~ 1	_	Colorless clear
IIb	~ 1	~ 100	Blue

to increase re-nucleation rate [30, 31]. The surfaces of MCD and NCD films generally get rougher, with highly faceted topography, typically $\sim 10\%$ of the film thickness, as the film thickness increases (Figs. 3b, c). In contrast to the growth process for MCD and NCD films, the UNCD film growth uses an argon (Ar) gas-rich chemistry (Ar (99%)/CH₄ (1%)), and no H_2 [2, 4]. As a result of the unique patented Ar gas-based gas chemistry, UNCD films provide a fundamentally different nanostructured PCD with grain sizes in the range 3–5 nm, the smallest diamond grain sizes compared to any MCD and NCD films. Figure 3d shows an SEM image of agglomerated UNCD grains, which can be seen only by HRTEM as shown in Figs. 3d, 33a,b of this review. In addition, UNCD films of any thickness exhibit roughness in the range 3-5 nm due to the high re-nucleation rate resulting from the unique Ar/CH₄ plasma chemistry

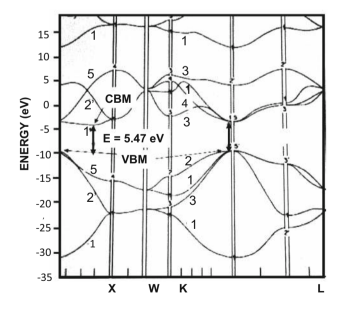


Figure 2 Energy-band structure of diamond calculated using the Discrete Variational Method (DVM) with Xo exchange scaling; CBM, Conduction Band Minimum; VBM, Valence Band Maximum, Fig. 2 in Reference [22].

[4, 32, 33], whereas NCD films turn into conventional MCD films after a certain thickness, due to grain size growth as a function of proportional film thickness increase.

In addition to PCD films, DLC films are also used for fabrication of MEMS devices. DLC films exhibit metastable amorphous carbon structure, involving a microcrystalline phase of diamond. AFM images of DLC films grown at different temperatures on glass are shown in Figs. 3e,f [34, 35]. DLC films involve sp^2 and sp^3 carbon bonds and are generally smooth and grown at low temperatures. Many DLC films are grown using a plasma-enhanced CVD (PECVD) process, ion beam sputter-deposition, or laser ablation of graphite solid target [3], all of them involving different gases mixtures. Some DLC films do not have hydrogen in their solid form (a-C or ta-C), while other DLC films have H atoms inserted in their structure (a-C:H or ta-C:H). Both a-C and a-C:H DLCbased films exhibit high sp^2/sp^3 C atoms bonds ratio. However, DLC films may exhibit high fraction of sp^3 bonding, when grown using enhanced plasma density sputtering or CVD processes. In order to distinguish them from sp^2 bonded DLC films, McKenzie suggested to name preferential sp^3 bonded DLC films as tetrahedral amorphous carbon (ta-C) films [34] (see also [35]), while the ta-C:H -based DLC films were distinguished as hydrogenated tetrahedral amorphous carbon by Weiler et al. [36].

Raman spectroscopy is one of the main characterization tools used to determine in a rapid analysis the diamond film characteristic as MCD, NCD or UNCD, as shown in Fig. 4. SCD and MCD films show a sharp, narrow/high intensity peak at 1332 cm⁻¹ in the Raman spectrum, with a very small Full Width at Half Maximum (FWHM) value as shown in Fig. 4a [37]. The D and G bands characteristic of polycrystalline graphite at around 1345 and 1560 cm⁻¹, and a low intensity band centered approximately at 1470 cm⁻¹ correlates with a tetrahedrally bonded

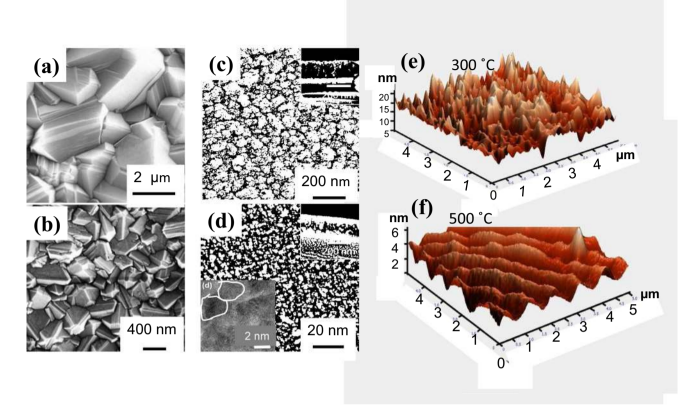


Figure 3 SEM images of PCD films grown on Si: a microcrystalline diamond (MCD/2–5 μ m grains) [318], b large grains (100–400 nm) nanocrystalline diamond (NCD) [318], c small grains (20–100 nm) nanocrystalline diamond (NCD) [318]; d ultrananocrystalline diamond (UNCD/ grain sizes of

diamond precursor [38]. The Raman spectrum of NCD, UNCD and DLC films are shown in Figs. 4b–d, respectively [39]. MCD has the highest diamond-type quality among PCD films, indicated by the high intensity of the sp^3 peak at 1332 cm⁻¹ and amorphous-like DLC film has the lowest diamond-like quality as indicated by the almost invisible sp^3 peak.

Table 3 shows the basic properties of various amorphous and crystalline forms of carbon provided in references [2, 36, 39–43].

Electrical conductivity of PCD

Surface electrical conduction

Undoped, high-quality diamond is one of the best insulator materials. However, diamond with H atoms-terminated surface exhibits relatively high

3-5 nm) [318]; diamond-like carbon (DLC) grown on glass at 300 °C (e) and 500 °C (f) respectively, showing that the surface roughness may depends on the growth temperature among other factors [35].

conductivity in air. The conductance found in the experiments is summarized in Fig. 5 [44, 45].

The electrical conductivity of H atoms-terminated diamond surface is induced by positive-charged carriers (holes) concentrated in a narrow surface region. Several proposed explanations are not widely accepted yet. The surface transfer doping mechanism proposed by Maier et al. [44] and Ristein, et al. [45], and Behravan [46] is in general agreement with experimental measurements of electrical conductivity on diamond surfaces. Figure 6 shows schematics of the basic concept of the surface transfer doping mechanism: Exposing a hydro-germinated surface to normal ambient air, a thin layer of adsorbed waterabout 1 nm thick—forms at the hydrogen-terminated surface. Assuming that this adsorbed water layer is neutral, i.e. in case it does not contain any acid- or base-forming impurities, small and equal densities of H_3O^+ and OH^- ions are formed:



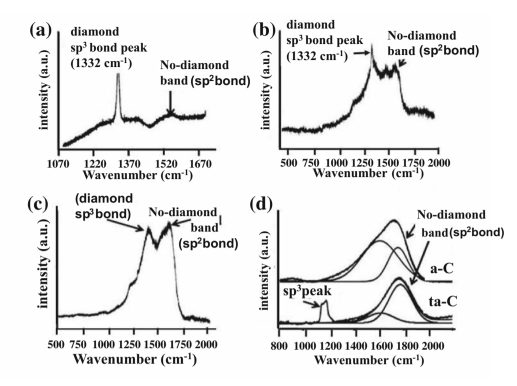


Figure 4 Raman spectra obtained with visible LASER with 532 nm wavelength: a microcrystalline diamond (MCD) [Auciello's group figure], b nanocrystalline diamond (NCD)

[33], **c** ultrananocrystalline diamond (UNCD) [33], **d** diamond-like carbon (DLC), including a-C and ta-C [36].

 Table 3 Properties of various forms of carbon

	Density (g/ cm ⁻³)	Hardness (Gpa)	Young's modulus (GPa)	<i>sp</i> ³ (%)	H (at%)	Gap (eV)	Surface roughness	
SCD	3.52	100	1050-1200	100	< 0.1	5.45		
MCD (grain size $\sim 0.5-10 \ \mu m$)	3.52	70–100	800-1200	~ 100	< 1	5.45	400 nm – 1 μm	
NCD (grain size 50–100 nm)		30–75	800–1020	> 50	< 1	2-4.7	50-100 nm	
UNCD (grain size $\sim 3-5$ nm)	3.50	88–98	916–980	95–98	< 1	5.4–5.65	20-40 nm	
DLC	ta-C (evaporated)	~ 3.00	2–5	757			3–3.5	5–100 nm
	ta-C (MSIB)	3.0	30–130		90 ± 5	< 9	0.51.5	5-100 nm
	ta-C:H	2.9	61		75	22-28		
	a-C:H (hard)	1.6-2.2	10-20	300	30-60	10-40	0.8 - 1.7	1-30 nm
	a-C:H (soft)	0.9-1.6	< 5	300	50-80	40-65	1.6–4	1-30 nm
Graphite	2.267	1–2	9.2–13	0		-0.04		
Glassy Carbon	1.3–1.55	2–3	35	~ 0		0.01		

$$2H_2O \leftrightarrow H_3O^+ + OH^- \tag{1}$$

Transferring valence electrons from the diamond valence band to the dissolved H_3O^+ ions, a thin sheet

of holes below the diamond surface and a layer of compensating negative OH⁻ ions in the adsorbed surface electrolyte are formed. A much higher surface conductivity arises, in case acid-forming molecules

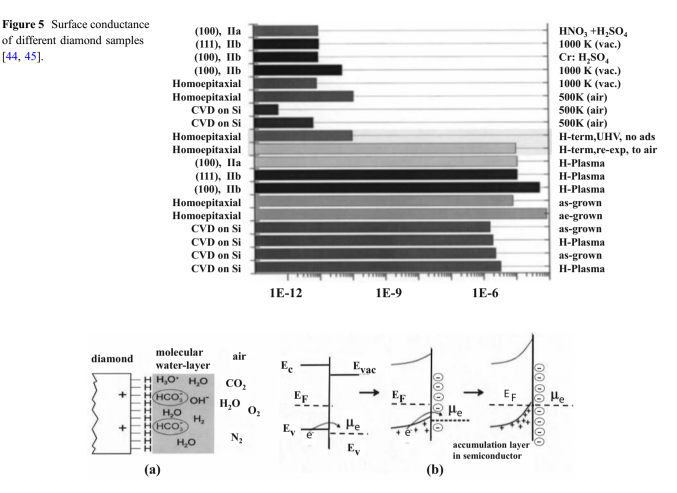


Figure 6 a Schematic of the diamond surface with adsorbed HCO_3 molecules, resulting from expsure to air, and inducing H-termionated diamond surface **b** evolution of the band bending

become dissolved in the electrolyte layer. The acid surface electrolyte layers may arise from the dissolution of atmospheric CO₂:

$$CO_2 + 2H_2O \leftrightarrow H_2CO_3 + H_2O \leftrightarrow H_3O^+ + HCO_3^-$$
(2)

In this case, a much higher sub-surface hole density is expected and the HCO_3^- ions in the electrolyte now take over the role of the anions. This is illustrated in Fig. 6a. Water vapor from the atmosphere reacts chemically with the diamond surface resulting in a catalysis process, whereby electrons are extracted from the diamond surface and inserted to a HCO_3 adsorbate on the diamond surface, contributing to electrical conductivity. The formation of holes on the diamond surface sketched in Fig. 6b. The electrochemical potential (Fermi energy) μ_e of the adsorbed electrolyte layer on the diamond surface is fixed by a redox level. Due to the surface of an adsorbate layer

during the electron transfer process at the diamond/ adsorbed HCO₃ surface layer intereface [46].

getting in contact with the diamond surface, electrons transfer from the adsorbed surface layer on to the diamond surface, resulting in the transition of the Fermi energy level of the solid to equal the Fermi energy level of the ambient film, similar to the case of a metal–semiconductor contact, with the difference that the adsorbed surface layer on the diamond surface replaces the metal electrode layer on the semiconductor. The accumulating holes (resulted from the electron transfer from the valence band) in the diamond near the surface and negative charges of HCO_3^- on the diamond surface result in band bending near the surface and thus a finite surface conductivity.

The maximum of the hydrogen-terminated diamond valence band coincides with one of the redox levels in the electrolyte layer. H_3O^+ ions therefore can become neutralized by the transfer of valence electrons to the liquid electrolyte. As a consequence, a high hole density is able to build up at the diamond surface. If the diamond surface is oxygen-terminated, huge thermal activation energy is required for the transfer doping. A p-type surface electrical conduction, therefore, does not arise on oxygen-terminated diamond.

The surface conductivity can be removed or changed by various surface treatments [33]: chemical cleaning of the surface with acids such as HF, HNO₃, HCl, H₂SO₄, CrO₃ and HClO₄, electrochemical cleaning, oxygen annealing, heating in ultra-high vacuum at 950 °C, heating in air 700 °C, oxygen plasma cleaning, N₂-Ar plasma and He-O₂ plasma.

Electrical conductivity of bulk diamond

The electrical resistivity of natural diamond is approximately 10^{14} – $10^{16} \Omega \cdot cm$ [47–50]. Owing to its high electrical resistivity, diamond is a good candidate for insulating layer material in MEMS/NEMS device. However, producing SCD on large area Si wafers, which may be required to produce low cost diamond-based MEMS/NEMS devices, may require substantial new R&D, involving several years in the future. Therefore, an alternative, for a more immediate implementation of diamond-based MEMS/ NEMS devices, is to develop processes for growing PCD films on large area Si wafers. PCD films grown by CVD processes can be micromachined to fabricate MEMS/NEMS devices. After appropriate surface treatment is done to remove the PCD film's surface electrical conductivity as described in Sect. 3.3.1, sometimes the as grown undoped PCD films show a resistivity of a few orders of magnitude lower than that in natural diamonds. The electrical conduction mechanism in undoped PCD films has been extensively investigated [51–64]. Two main mechanisms have been proposed for diamond films, namely fieldactivated (FA) emission [52, 54] and space-chargelimited (SCL) transport [53]. The former contribution can be represented by the Frenkel-Poole equation [58].

$$G(V) = G_0 \exp(\alpha V) \tag{3}$$

with

$$\alpha = q \Big/ 2kT dN_t^{1/3} \tag{4}$$

G(V) is the electrical conductance of diamond film, controlled by the drop of electrical potential across the diamond layer. N_t is the number of traps per unit

volume in the diamond layer, such that the coulombic potentials of traps overlap, and d is the diamond thickness; T is the absolute temperature and k is the Boltzmann constant. Research demonstrated that FA transport is dominant in diamond films containing large number of defects per unit volume, whereas diamond layers with large grains exhibit quality and performance of SCD. The high field conductance associated with SCL current can be represented by Eq. 5 below [58].

$$G(V) = c_1 V^l \tag{5}$$

where c_1 is an experimentally determined coefficient and *l* is an exponential in the range 0.5–3.5 [52]. A typical Current–Voltage (*I–V*) measurement on undoped PCD films, with dual side contact, made to the top and bottom surfaces of the diamond film, is shown in Fig. 7 [52]. For low voltages (< 10 V), *I–V* is linear and symmetric for both gold and indium contact layers. At high voltages, the current increases more rapidly than linearly. The nonlinear behavior at high voltages was explained by FA emission.

The FA emission modeling above considers the electrical conduction of highly defective grain boundary regions and grains at the same time as a whole. A more complete model proposed by Cesare et al. [58] separate the electrical conduction of grain boundaries from the grains. Electrical conduction through grains is assumed to follow the FA emission modeling, but grain boundary regions don't. The modeling involved also consideration of a rectification process between the metal electrode layer (contact) and the bulk of the diamond film, instead of using the ohmic contact concept, because contacts

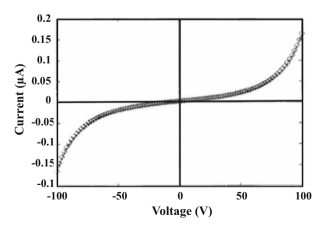


Figure 7 *I–V* characteristics of an undoped PCD sample with gold contacts top and bottom. [53].

with rectification usually happens on undoped diamond. The model considers two conductance phenomena, *G* and *g*, in parallel, characteristic of a combination of bulk crystal and highly defective grain boundaries, respectively, The model indicates that an area with a diode *D* characteristic exist in series with an area with electrical conductance *G*, representing a rectifying contact between at the metal electrode/diamond interface and the bulk region of the diamond layer. The model is based also on the assumption that an ohmic contact exists between the metal electrode and the highly defect grain boundaries of the diamond layer, all represented by Eqs. (6) and (7)

$$I = G(V_G)V_G + g_0 V_a \tag{6}$$

$$G(V_G)V_G = I_0\{\exp[q(V_a - V_G)/nkT] - 1\}$$
(7)

I is the total current when a voltage V_a is applied. V_G is the potential drop across the conductance *G*, *n* and I_0 are the quality factor and reverse saturation current of the diode, respectively, and *q* is the electron charge. The electrical conductance of grain boundaries is assumed to be a constant g_0 . Conductance of grains $G(V_G)$ can be expressed using the FA emission model given by Eqs. (3) and (4). This electrical model can reproduce the experimental *J*– *V* curves with adequate accuracy, as shown in Fig. 8.

The electrical conduction of undoped PCD can be attributed to the impurities or traps in diamond grains and grain boundaries. After comparing the quality of diamond grown under different CH_4/H_2 , revealed by SEM and Raman analysis, Kulkarni et al. attributed the low resistivity on the order of 10–10⁶ Ω cm to the grain boundaries structure [56, 57]. Lee produced Cu electrode layers, via electroplating, on diamond film surfaces, showing that the main electrical conduction path in undoped PCD films are the grain boundaries within the films [61]. The electrical conduction mechanism, due to impurities [62, 63], had also been investigated by examining the relationship between the material's temperature and the electrical current through the diamond film. The activation energy can be calculated by using the Eq. (8)

$$\sigma = \sigma_0 \exp(-E_a/kT) \tag{8}$$

where σ_0 is a pre-exponential constant and E_a is the donor/acceptor activation energy, which is related to the transition of electrical carriers from the trap to the

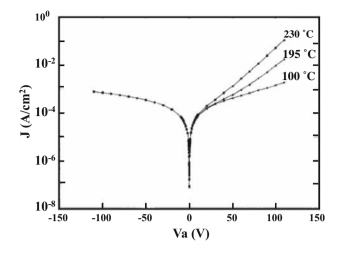


Figure 8 *J–V* curves resulting from calculations using the model described in Sect. 3.3.2 [58].

conduction/valence band levels, respectively. Thus, the type of impurities will be known once the activation energy is derived. Usually, the presence of dopant B atoms in the diamond lattice correlate with electrical activation energy in the range 0.3-0.4 eV. On the other hand, impurity N atoms in the diamond lattice, correlate with activation energies in the range1.4-1.6 eV. The electrical conductivity vs 1/T (temperature), shown in Fig. 9 [55], suggests that the electrically active diamond induced by B and N atoms incorporation into the diamond lattice is correlated with the film grown along (111) and (100) directions, respectively. Wang, et al.[60] also reported activation energy of 1.68 eV, for electrical conduction, attributed to Si atoms trapped at a vacancy in the diamond lattice. He speculated that the possible source of B atoms contamination in their diamond films could arise from the Si wafer, heavily doped with B atoms. Stoner et al. found that H atoms trapped in the film can also be an origin for electrical conduction of undoped PCD films with activation energies less than 0.6 eV [64], although they didn't remove the surface conduction layer, making their experimental results still intriguing. Stoner et al. reported that the initial room temperature electrical conductivity can change by 5 orders of magnitude, even though the relative graphite, or non-diamond structured, revealed by SEM, Raman and photoluminescence analyses, is nearly identical. When the temperature is > 500 K, the activation energy as approximately 1.1 eV. It was speculated that this transition, above 500 K, occurs because the H atoms in the lattice becomes mobile and then no longer



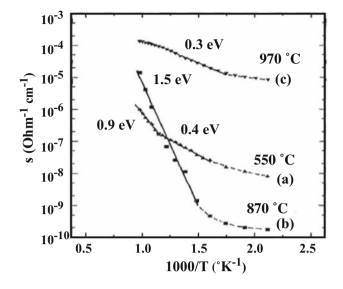


Figure 9 Electrical conductivity σ plotted versus 1/T for three diamond films grown under different temperatures [55].

occupies the various trapping sites. The real cause for low resistivity of undoped PCD films still needs further study.

Piezoresistivity of PCD films

The piezo-resistive effect was first discovered in 1856 by Lord Kelvin using copper and iron wires. The discovery of piezo-resistance in Si and Ge materials at Bell Laboratories in 1954 by Smith [65], is considered one of the milestones that fueled the growth of the MEMS industry. Smith found that the semiconductors exhibit piezo-resistance coefficients much higher than those of metals, which stimulated increased interest in the investigation of the effects of stress on other semiconductors and in the application of this effect to MEMS sensors.

Piezo-resistance is a change in resistivity of the material under the influence of an externally applied stress or strain. The piezo-resistivity of a material is related to the change of electrical resistivity as strain is applied, and is quantified by the gauge factor (GF). The GF of a material is defined as the fractional change of resistance ($\Delta R/R$) per unit strain ε , which is the figure of merit for piezo-resistive material, expressed mathematically by Eq. 9 below.

$$GF = (\Delta R/R)/\varepsilon \tag{9}$$

Typical GFs for various materials in the longitudinal direction are shown in Table 4 [66]. The reported GFs of PCD, in the literature, are given in

Material	Description	Gauge Factor (GF)
Metals	Composition	
Nickle/Copper	45% Ni, 55% Cu	2.0-2.1
Copper	100%	2.6
Silver	100%	2.9
Platinum/Tungsten	92% Pt, 8% W	4.0
Platinum	100%	6.1
Nickel	100%	-12.1
Material	Description	Gauge Factor (GF)
Semiconductors	Types	
Single Crystal Silicon	p-type	100–175
	n-type	-133
Poly-Silicon	p-type	15-30
	n-type	-30
Germanium	p-type	48.7-101.5
	n-type	- 147 to - 157
Poly-Germanium	p-type	30
	n-type	-30 to -40
Silicon Carbide	n-type	- 55 to - 994

 Table 4 Gauge factors for various materials in the longitudinal direction [66]

Table 5 and they were found to increase from 5 to 100, with increase in resistivity, grain size and operation temperature.

The piezo-resistive property of a semiconductor such as doped diamond, under uniaxial stress, can be explained by the valence-bands split-off model, as shown in Fig. 10.

The strain will change the concentration of heavy and light holes, so the strain can change the electrical conductivity of the p-type semiconductor. The GF for single crystalline doped diamond is given by the formula 10 below [68]

$$GF = -\frac{1}{\varepsilon} \frac{A + E_e C}{kT + A + E_e C} \approx \frac{1}{\varepsilon} \frac{A + E_e C}{kT}$$

$$C = \left(\frac{m_l^{1/2} - m_h^{1/2}}{kT} \right) / \left(\frac{m_l^{1/2} + m_h^{1/2}}{kT} \right) \text{ tensile strain}$$

$$C = \left(\frac{m_h^{1/2} - m_l^{1/2}}{kT} \right) / \left(\frac{m_l^{1/2} + m_h^{1/2}}{kT} \right) \text{ compressive strain}$$
(10)

where *A* and E_e are labeled in Fig. 10. The m_h and m_l coefficxients are the effective mass of heavy- and light band holes. The suffix '*h*' and '*l*' represent heavy and light holes, respectively. *C* represents the relative difference between the effective mass of heavy and light holes, in other words, they also represent the relative difference between the effective mobility of

GF	$\rho \; (\Omega {\cdot} cm)$	Boron Concentration (cm ⁻³)	Substrate	Deposition	Doping	References
6		1.2×10^{16}	Diamond/SiO ₂	HFCVD		[203]
5.4	4		Undoped carbon	MPCVD	BCl ₃	[102]
116		2.5×10^{18}	Si	DC plasma CVD	B ₂ O ₃ powder	[206]
6–25	5-30		SiO ₂	HFCVD	boron powder	[67]
8	0.2		Si	MPCVD	Boron compound solid wafer	[219]
10	2.2		Si	MPCVD	Boron compound solid wafer	[219]
67	225		Si	MPCVD	Boron compound solid wafer	[219]
100	300		Si	MPCVD	Boron compound solid wafer	[219]
1000	300		Undoped PCD	MPCVD	B_2H_6	[208] [209]
7–9	0.1		Si	MPCVD	B(CH ₃) ₃ in ethanol	[210]
70–75	100		Si	MPCVD	B(CH ₃) ₃ in ethanol	[210]
0.1	1000		Si	MPCVD	B(CH ₃) ₃ in ethanol	[210]
30–35	10^{6}		Si	MPCVD	B(CH ₃) ₃ in ethanol	[210]
690			Si	MPCVD	B(CH ₃) ₃ in ethanol	[211]
283*	0.27		Undoped PCD	MPCVD	boron powder	[212]
4000**	0.1					
6.7/8.9*		$\sim 10^{20}$	Undoped PCD	MPCVD		[213]
25**		~ 1020	Undoped PCD	MPCVD		[213]
33	3.15		SiO ₂	HFCVD	H ₃ BO ₃ in methanol and acetone	[215]
50	2.65		SiO ₂	HFCVD	H ₃ BO ₃ in methanol and acetone	[215]
40	28		SiO ₂	MPCVD	B(CH ₃) ₃	[72]
8 ± 0.5	0.01		SiO ₂	MPCVD	B(CH ₃) ₃	[217]

Table 5 Gauge factor of PCD reported by different groups

*inter-grain piezo-resistor; *intragrain piezo-resistor

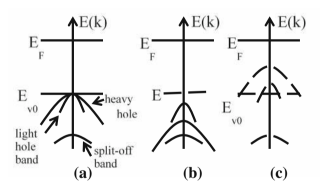


Figure 10 Simplified Schematic of valence bands split-off model a no stress, (b) under tensile stress, and (c) under compressive stress.

heavy and light holes. For diamond, C = -0.89 under tensile strain and C = 0.89 under compressive strain. k is the Boltzmann constant, T (K) is the absolute temperature. The great difference of the effective mass between the heavy and light holes in diamond is one of the main factors causing the excellent piezo-resistive effect of diamond [68], as indicated by Eq. (10). Since *GF* is positive under

tensile, it indicates the resistivity increase under tensile stress. Likewise, the resistivity decreases under compressive stress due to the negative value of *GF*.

Generally, polycrystalline materials are formed by crystalline grains interconnected by grain boundaries. In the case of diamond, grains are made of single crystals with the diamond lattice. The grain boundaries (GBs) involve C atoms in a disordered structure with a large number of defects and dangling bonds, acting as trapping states and/or segregation sites [69], for trapping of holes, creating an electric potential barrier at the boundary and depletion regions into the grains. The diamond film may be modeled in terms of grains of length L_g with boundary width δ and depletion width w_d [70]. The change in resistivity per unit strain of the polycrystalline materials can be calculated using Eq. 11 [71]:

$$\frac{\Delta\rho}{\rho}\frac{1}{\varepsilon} = \frac{L_g - (2w_d + \delta)}{L_g}\frac{\Delta\rho_g}{\rho_g}\frac{1}{\varepsilon} + \frac{(2w_d + \delta)}{L_g}\frac{\Delta\rho_b}{\rho_b}\frac{1}{\varepsilon}$$
(11)

where ρ_g and ρ_b are the resistivity of grain and grain



boundary, respectively. Research suggests that the piezo-resistive effect indued by the GBs is much lower than the one induced by the crystalline grains. The random orientations of the GBs also lower the overall piezo-resistive effect of grains, which correlates with the reported low GF of polycrystalline materials compared to single crystal counterparts. The effect of GBs on the gauge factor was investigated theoretically [68] and described by the Eq. 12 below, assuming there is no gauge factor for GBs:

$$GF \approx \left(\frac{1}{1+\alpha}\right) GF_g$$
 (12)

where α is the GBs / grains resistances ratio through the electrical conduction path when the film exhibits no strain, and GF_g is the gauge factor of single crystalline material. The calculations using Eq. 12 explains why polycrystalline films, with high resistivity, exhibit larger *GF*. A more explicit equation of GF for a polycrystalline material is given by Eq. 13 [68].

$$GF = (M - F)GF_g + F(D - C)\frac{1}{kT}\frac{dE_e}{d\varepsilon}$$
(13)

D and C constants relating to the difference of the effective mass of holes. For diamond films, D has values of -1 and 1 under tensile and compressive stress, respectively. *M* and *F* are two constants related to scattering of electronic carriers in grain-boundaries. For a very small strain ($\sim 10^{-6}$), M is equal to 1 and 0 < F < 1. The last term on the right of Eq. (13) is negative, which indicates that GF is smaller than GF_{g} , correlating with the observation that the GF of polycrystalline materials is smaller than that of single crystalline materials. Equation (13) also indicates that if there is an excellent piezo-resistive effect in the crystalline materials, a more significant piezo-resistive effect will exist in the corresponding polycrystalline material, which support the concept that PCD is a good candidate for the application of MEMS piezo-resistive sensors.

Polycrystalline diamond (PCD) film technology

For the implementation of diamond as a key material enabling a new generation of MEMS/NEMS devices, successful growth of PCD films, doping, patterning and metallization processes must be achieved.

Chemical vapor deposition of PCD films

CVD diamond films have been grown on substrates of different materials. Monocrystalline diamond substrates induce homoepitaxial crystalline films, whereas non-diamond and PCD substrates result in the growth of PCD films. For non-diamond substrates, a nucleation procedure is necessary to produce the diamond precursors, namely: (1) surface abrasion with diamond particles to insert them on the substrate's surface [72–74], (2) sonication of diamond powder loaded solution [75], (3) bias enhanced nucleation (BEN) [76–78], where C = ions and $CHx^{=}$ (x = 1, 2, 3) from a plasma, created in front of the substrate surface, are accelerated, with very low energy, towards the surface, getting implanted on the surface and inducing the seeding process, (4) spinning of diamond-powder-loaded photoresist (DPR) [79], and (5) spraying of diamond-loaded fluids [80, 81] on the substrate surface. All substrate's surfaces seeding processes, described above, are applied to grown diamond films by different methods, as summarized in Table 6 [82]. Most of the previous seeding methods, described above, focused on seeding of bare Si wafer's surface or a tungsten surface and very few have reported diamond seeding on a dielectric surface. For the electrical application in microsystem, it is common to insulate a conductive film using dielectric layers (SiO₂ or Si_3N_4) and make connection through contact holes or vias. Thus, having the ability to achieve nondestructive, controllable, uniform and high-density diamond seeding on dielectric layer is an important step for PCD filmbased MEMS sensors.

Currently, PCD films are grown using various CVD methods, including microwave plasma CVD (MPCVD), hot filament CVD (HFCVD), radio frequency (RF) CVD (RFCVD) and dc-arc jet CVD (DCJCVD), as shown in Table 7 [82]. However, MPCVD is the most widely used technique for electronic and optical applications of diamond films, because of its high efficacy to produce high quality films on large substrate area, less contamination and good controllability of the growth process. The first PCD films consisted of randomly oriented grains interconnected by grain boundaries. The search for polycrystalline diamond films with improved electronic properties lead the development of deposition techniques for highly oriented films (Table 8).

Seeding Method	Sonication/abrasion	Ben	DPR	Spray/paint	DPL with spin
Seeding Density (cm ⁻²)	Up to 10 ¹⁰	Up to 10 ¹¹	$\sim 10^8$	Up to 10 ¹¹	10 ⁸ -10 ¹¹
Substrate Selectivity	Most dielectric & metal	Conductive Si or Metal	Most dielectric & metal	Most dielectric & metal	Hydrophilic surface (SiO ₂ & Si ₃ N ₄)
Substrate Surface Effects	Scratch surface, not good for thin film	No damage	No damage	No damage	No damage
Uniformity & Controllability	Not uniform	Uniform & repeatable on whole wafer	Uniform & repeatable on whole wafer	Not uniform	Uniform & repeatable on whole wafer

Table 6 Comparison of different seeding methods on seeding density, substrate material and surface morphology [83]

Table 7 Comparison of different CVD methods for growth of PCD films [82]

Methods	HFCVD	MPCVD	DC-arc Jet CVD	Combustion Synthesis	RFCVD
Growth Rate (µm/ hr)	0.1–10	0.1–10	30–150	4-40	< 0.1
Subst. Temp. (°C)	300-1000	300-1200	800-1000	600-1400	700-1200
Growth Area (cm ²)	5–900	5–100	< 2	< 3	100
Advantages	Simple, large area	Quality, stability	High rate, good quality	Simple, high rate	High quality
Disadvantages	Contaminations, fragile filament	Rate	Contaminations, small area	Small area	Low rate, expensive

Bachmann *et. al.* (in 1991) described the diamond growth region with the C–H–O gas phase diagram, as shown in Fig. 11 [83]. The C–H–O diagram gives a clear picture on where to search for diamond. It is apparent that MCD and NCD films can be grown using a wide range of gases mixture flown inside evacuated chambers with the gas mixtures including hydrogen (H₂), methane (CH₄), ethane (C₂H₆), ethylene (C₂H₄), acetylene (C₂H₂), and carbon dioxide (CO₂). The diagram is independent of deposition technique but not independent of temperature. One of the general trends revealed by Bachmann's review is the improvement in MCD and NCD film quality at the expense of growth rate with the decrease of carbon content and increase in oxygen content.

Bachmann et al.correlated the temperature of the CVD gas phase with the growth rate and film growth method, as shown in Fig. 12. The increase in deposition rate with gas temperature is generally

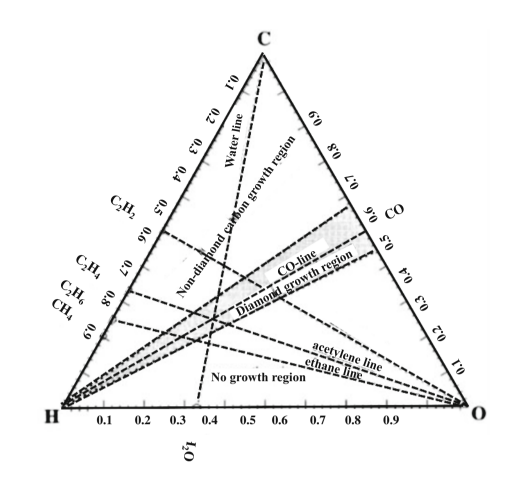
attributed to a more efficient supply of the diamond growth species. Investigations of diamond film's growth mechanisms show that methyl and acetylene are the main growth species for MCD and NCD films [84]. Besides gas composition and temperature, gas pressure and substrate temperature control the MCD and NCD film's growth process. Successful growth of MCD and NCD films were reported for substrate temperatures in the range of 400-1350 °C where poor quality sets the low temperature limit and absence of growth sets the high one. Bachmann et al. observed that the "diamond domain" narrows as substrate temperature increases. Gas pressures in the range of 1-200 Torr were used to grow MCD and NCD films, depending on the substrate temperature, gas temperature and gas composition.

Currently, designers of CVD diamond film growth reactors are pressured to deliver cheaper and better diamond films quality on large areas. The major

Table 8 Various diamond polishing techniques [192]

Technique Bulk Proce Temp	Bulk Processing Temperature	Polishing Shape mechanism limitati	Shape limitations	Shape Si A' limitations limitations	Special requirements	Set-up	Equipment cost	Equipment Large area cost processing cost	Processing Reported Parametric time(per roughness freedom cm ¹)	Reported roughness	Parametric freedom	Surface change (contamination)
Mechanical hipping	Room	Abrasive wear Only planar surfaces	Only planar surfaces	No limit	None	Rigid and geometry sensitive	Low	Low	Few days	210 nm	Small	Yes
Thermo- mechanical	D° 006-008	Graphitization, Only planar dirTusion - surfaces	Only planar surfaces	Plate size	Need environ- mental control	Rigid and geometry sensitive	Low	Low	Tens of minutes	5.5 run	Restricted	Yes
CAMPF	> wc	Oxidation	Only planar surfaces	Plate size	None	Simple	Low	Low	A few hours	49 nm	Restricted	Yes
Laser	Room	Etching, evaporation	Non-planar surfaces nossible	No limit	Need scanning of the samples	Simple	High	Medium	Few Seconds	500 m	Large	Yes
Ion beam	Room	Sputtering etching	Non-planar surfaces possible	Beam size	NeeJ high vacuum	Complex	High	High	Tens of hours	Mint	Restricted	Yes
RN;	700 = 0	Sputtering	Non-planar surfaces possible	Plasma size	NeeJ high vacuum	Higid	High	Medium	Tens of minutes	71 nm	Large	Yes

Figure 11 Phase diagram by Bachmann et al. indicating the possible gas combinations in reported successful PCD film growth experiments [83].



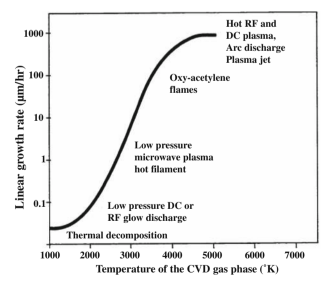


Figure 12 Growth rate for different CVD methods correlate with the temperature of the gas mixture needed for growth [83].

challenges they face to apply PCD films to fabrication of MEMS devices involve producing large area diamond films with uniform thickness and micro- or nanostructure across the whole area, low temperature deposition and small residual stress in the films.

Doping of PCD films

Accurate control of the doping level and resistivity is particularly important for electrical applications of PCD thin film in sensors. Since the 2000 K temperature necessary for effective diffusion of dopants in diamond is too high [85], diamond film doping is performed either during film growth or by subsequent ion-implantation. Boron (B), Aluminum (Al), Phosphorous (P), Lithium (Li) and Nitrogen (N) atoms have been tested as dopants for SCD and PCD films [86]. Currently, B is the most successfully used dopant to fabricate diamond film-based electronic devices. Different B-containing compounds, typically used for p-type doping, are pure B-atomsbased powder [87], B-trioxide (B₂O₃) [88], diborane (B_2H_6) [89, 90] and trimethyl boron $(B(CH_3)_3, TMB)$ [91]. Although B^+ ion implantation has been used for diamond doping [92, 93], its effectiveness has been





limited by the implant damages and very low diffusion ability of B atoms in PCD films [94]. Use of TMB to produce B atoms doping of diamond films is currently a preferred source of B dopants because the TMB gas, diluted in hydrogen, is safer to use and more controllable than other B-based compounds [91]. IR measurements confirmed that B atoms occupy substitution sites in the diamond lattice [95]. Figure 13 shows the activation energy and hole concentrations of B-doped PCD films under different doping levels and temperatures [96]. The B activation energy in PCD films was found to decrease with doping, starting at 0.37 eV [97]. This decrease has been attributed to the appearance of an impurity band that increases in width until the onset of metallic conduction at B atoms concentrations greater than 10^{21} cm⁻³ [98].

A convenient equation used to relate the activation energy to the doping concentration is shown below. This means that for doping concentrations N_A greater than 1.7×10^{20} cm⁻³ the activation energy E_a becomes zero, correlated with experimental measurements [99].

$$E_a = 0.37 - 6.7 \times 10^{-8} N_A^{1/3} \text{ eV}$$
(14)

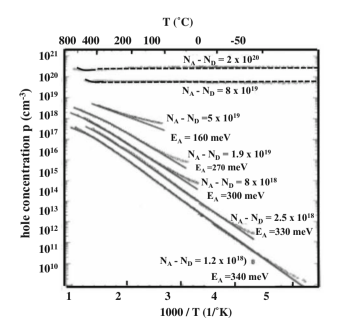


Figure 13 Activation energy and hole concentrations of boron doped PCD films under different doping levels and temperatures [98].

Patterning of PCD films for MEMS fabrication

For integration of PCD films into Si-based MEMS/ NEMS devices, patterning of PCD films, based on Si processing techniques, are required. As the wet etching of PCD films is impractical, since diamond is resistant to all chemical etchants, the commonly used patterning techniques are in-situ pattering and selective dry etching. Laser beam [100] and synchrotron excited radiation [101] were also reported as an optional way to etch PCD films.

On the other hand, selective growth of PCD films on previously patterned structures can be used to produce patterned diamond film-based structures, which can be achieved by selective nucleation of diamond or by masking the areas where growth is not desirable. SiO₂ was successfully used, as a masking layer, by Masood et al. [102], Roppel et al. [103] and Davidson et al. [104]. Patterned amorphous Si masking method was also reported [105]. Hirabayashi et al. first nucleated the Si substrate by ultrasonic treatment, then used a photoresist mask for etching Si to 60–70 nm, using an Ar+ ion beam [106], or reactive ion etching [105]. A simple selective diamond nucleation technique, based on spinning a photoresist/diamond powder pre-mixed layer In the substrate surface, followed by lithographically patterning, was developed by Aslam's group at Michigan State University [102].

However, in all the in-situ nucleation techniques, the lateral growth at the pattern edges and, in some cases, growth on unwanted areas result in serious problems for fabrication of small feature sizes for NEMS devices. Consequently, selective dry etching is an excellent choice for patterning of PCD films. In this sense, selective etching of PCD films, using SiO₂ or Si₃N₄ as masks, was performed at atmospheric pressure, in oxygen environment at 700 °C, in a rapid thermal processor [102]. Dry etching technique using conventional dry etching tools, include: (1) ion beam etching (IBE), (2) reactive ion etching (RIE), (3) electron cyclotron resonance etching (ECR) and (4) inductively coupled plasma etching (ICP), have been intensively investigated and developed in recent vears.

Metal films for electrical contacts on diamond films

Metal contacts to diamond have been investigated extensively, due to their importance in the electrical characterization of diamond films. Two types of metals have been used extensively, namely: carbide and non-carbide forming metals. Carbide forming metals (e.g., Ti, Mo and Ta) [107–109] provide ohmic contacts upon annealing. Low contact resistivities $(\sim 10^{-7} \,\Omega \cdot \text{cm})$ were achieved for Al/Si-contacts [110] on diamond, after annealing at 450 °C. The low contact resistivity has been attributed to the formation of SiC on the Si substrate surface in contact with the metal electrode surface. Research suggest that the SiC layer contain defects, which lower the barrier height or enhances electrical carriers tunneling, or both. It is possible that annealing may increase the local disorder via introduction of point defects. The model to explain the ohmic contact resulting from carbide formation is shown in Fig. 14 [47, 111]. There is a barrier at the metal / diamond interface before annealing (Fig. 14a). The carbide formation produces point defects at the surface of the diamond. These defects can either decrease the width of the depletion layer, and therefore enhance the probability of tunneling for the electrical carriers (Fig. 14b) or reduce the effective barrier height (Fig. 14c). Although annealing helps the formation of ohmic contact, Ti can diffuse to the Au/ambient interface at high temperature [112–116]. Thus, Ti/Mo/Au, TiWN/Au or Ti/Pt/Au stacked layers are used to prevent diffusion. Ohmic contacts were observed after heavy B atoms doping either in-situ, during diamond film growth or implanted after film growth, regardless of the metal being used [117, 118]. Heavy doping is regarded as a process that narrows the width of the depletion region at the metal-diamond interface, increasing the electronic carries tunneling current. By using heavily doped PCD film as an interlayer between diamond and metal, the contact resistance is reduced to one third of the original value [119].

Figure 15 shows curves of contact resistivities, measured experimentally, for various electrode materials, by different groups after annealing [120]. Tunneling theory was used to produce the lines in Fig. 15, for comparison with experimental data (shown by dots, triangles, squares), which shows tunneling theory does not agree well with experimental data for high doping levels. The data in

Fig. 15 indicates that the use of single barrier height, after annealing, to model the doping dependence of the specific contact resistivity, is not reliable, and/or there is another neglected conduction mechanism, not accounted for in the modeling, such as the impurity band electrical conduction.

In addition to the work function and type of metal, surface treatments on diamond also contribute substantially to changing the Schottky barrier height, resulting in different contact resistivity. A comparison of Schottky barrier heights data, as measured on clean and oxygen-terminated diamond surfaces is given by Fig. 16. Obviously, oxygen-terminated surfaces lead to larger Schottky barrier heights. Mori also found that if the diamond surface is oxygen-free, the I-V behavior depends on the metal work function and its electronegativity. If oxygen is on the surface, this dependence vanishes [121]. This is in agreement with another group work, which also found that the metal-diamond interfaces, where diamond is H-terminated with negative electron affinity, lowers the Schottky barrier heights [122].

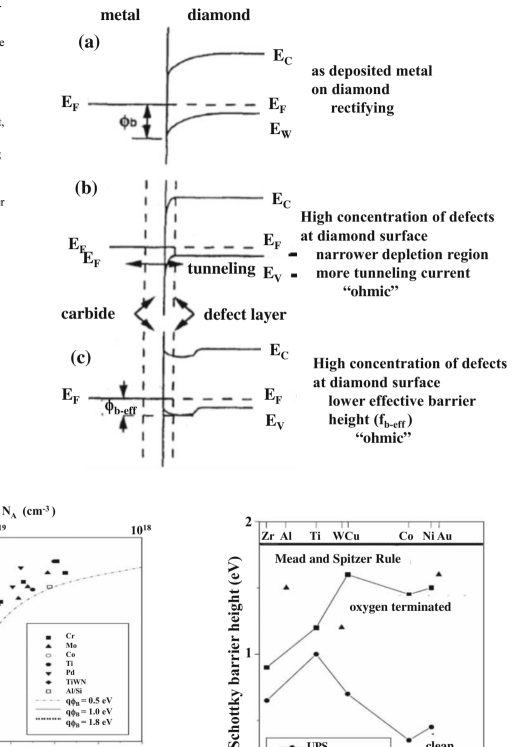
PCD films-based MEMS technologies

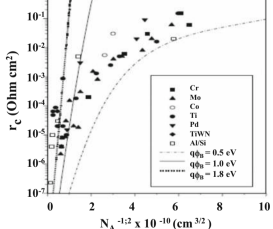
Dry etching of crystalline and polycrystalline diamond films

Four major methods of dry etching SCD and PCD films have been reported in the last thirty years, including ion beam etching (IBE) [123, 124], reactive ion etching (RIE) [124-145] electron cyclotron resonance etching (ECR) [146-153] and induced couple plasma etching (ICP) [154-156] Generally, O₂ has been widely used to etch diamond by generating volatile CO and CO₂ molecules [157], whereas diamond columns were found to stand on the etched diamond surface. Though these diamond columns are an excellent candidate for an electron field emitter [128, 130–132, 141, 143–145], it is not desirable for PCD films etching, as it hinders the removal of PCD layers to produce MEMS structures not requiring columns. In order to suppress and eliminate the diamond columns, a small additional amount of CF4 or SF_6 is introduced into the O_2 plasma, thereby sacrificing the etching selectivity of the Al or SiO₂ masking layer to the diamond film [126, 127, 129, 134, 135, 139]. However, the plasma environment with the presence of CF₄ or SF₆ also



Figure 14 Band diagrams of metal-diamond interfaces: **a** for as-grown metal electrode layer on diamond, b high concentration of electrically active defects, created via theformation of the carbide layer and/or ion bombardment, wich reduces the depletion width and increases tunneling current, **c** electrically active defects created as in **b** decrease the effective barrier height [111].





10¹⁹

10²¹ 10²⁰

10⁰

Figure 15 Experimentally determined contact resistivities for various electrodes materials, from different groups, following the annealing process. The tunneling theory does not agree with experimental data well for high impurity levels [120].

Figure 16 Comparison of Schottky barrier heights as measured on clean and oxygen-terminated diamond surfaces [120].

Metal Work-function (eV)

Electrical measure

4.5

-UPS

-UPS

٠

0

4.0

clean

5.5

5.0

etches Si, SiO₂ or Si₃N₄ very fast. Such damage is inevitable when doing etching of PCD films, mainly due to the existence of discontinuous areas in the PCD films at the last stage of PCD films' etching. Such discontinuity can be attributed to: (i) discontinuous areas in the PCD film near the interface between the PCD layer and the underlying layer in the beginning of the PCD film growth; (ii) surface roughness of as-grown PCD films; (iii) preferential etching of PCD films along grain boundaries or in certain facet orientation. Considering the above issues in etching of PCD films, a three-step etching scheme was developed, using a sequential RIE process in CF₄, O₂ and H₂ plasmas, to remove PCD film from areas required to be removed very efficiently, and also achieve a medium-smooth etched surface [158], as shown in Fig. 17. This result is very important to the application of PCD film to produce multilayer diamond-based MEMS structures.

Residual stress in diamond films

The stress produced in SCD and PCD films during the growth process affects strongly the physical properties of the grown films. The stress induced in the SCD and PCD films may result in peeling of the film from the substrate due to compressive strain. Therefore, it is critical to understand the underlying mechanism (s) that produce residual stresses in diamond films described here. Two main components of residual stress have been identified in SCD and PCD films, namely: (1) thermal stress, resulting from cooling the films from the temperature at which they are grown to room temperature; this thermal stress is

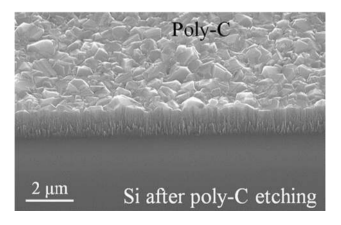


Figure 17 Dry etching of PCD film with medium smooth etched surface using a three-step etching scheme, i.e. a sequential RIE etching in CF_4 , O_2 and H_2 plasmas [159].

due to the difference between the thermal expansion coefficients between the diamond films and the substrates; (2) intrinsic stress, resulting from the film growth process; this intrinsic stress is related to the non-diamond phase at the grain boundaries of the PCD films and to structural defects, such as impurities, micro twins, and dislocations.

Techniques used to measure residual stress in diamond films include: (a) measurement of substrate curvature [159-163] analysis of X-ray diffraction (XRD) diamond peaks [161, 164-172] and measurements of peaks obtained in Raman analysis of the diamond films [161, 162, 164, 167-169, 173-178]. The measurement of the substrate curvature involves sending a focused laser beam onto the integrated diamond film/substrate structure and detecting the position of reflected beams on a screen; this procedure provides the information to determine the curvature of the wafer, produced by the stress induced by the diamond film grown on the substrate surface. using the Stoney's equation. XRD and Raman analvsis provide indirect ways to measure the diamond residual film stress when the integrated film/substrate curvature cannot be measured.

Raman spectroscopy provides a very simple and rapid way to measure the stress in SCD and PCD films, via characterization of the diamond peak located at 1332 cm⁻¹ in the Raman spectrum. In this sense, the Raman diamond peak is shifted from the 1332 cm⁻¹ position, as the stress in the diamond films distort their crystalline structure.

In XRD analysis, the shift of the diamond peak, from the position characteristic of unstressed diamond, provides also the means to measure the diamond film stress. However, the XRD-based measurement method, to determine diamond film stress, is statistically better that the Raman-based technique, because XRD analysis enables the evaluation of average stress on larger films' areas than that provided by the smaller area focused micro-Raman spectroscopy [166]. In addition, due to the general transparency of diamond films to X-rays, the XRD technique provides information about the stress of the films through the whole film thickness.

Measurements of stresses in diamond films have shown that the intrinsic stress depends on key parameters used for film growth, namely: CH_4 gas flow fraction, substrate temperature during diamond film growth [159] and crystalline orientation of the diamond films [179]. In undoped PCD films, the non-



diamond carbon phase, generally observed in grain boundaries, and H atoms content, have been considered as major contributors to inducing compressive stress in the diamond films [160, 175]. In addition, for PCD films, mechanical relaxation in the structure of grain boundaries is considered as a major contributor to tensile intrinsic stress [169]. For B-doped PCD films, prior research [173] showed that tensile stress is enhanced with increasing B atoms concentration in the diamond film. The tensile stress is attributed to grain boundary relaxation, due to the short-range B atoms-interatomic distance, resulting in B atoms concentration in the films' lattice below 0.3 at. %. On the other hand, for diamond films with high B-atoms concentration (> 0.3 at.%), the internal stress is attributed mainly to high defect density, induced by blocking of sliding processes in grain boundaries. The understanding of the mechanism (s) responsible for residual stress in diamond films, produced by the CVD process, is fragmented, with significant disagreement among various research groups, in relation to the mechanism (s) controlling the type and magnitude of the stress, even for similar film's growth conditions. The disparity in values of measured stress may be attributed to the stress measurement methods [161], which generally exhibit limitations to be considered.

Many applications of diamond coatings are strongly limited due to the poor adhesion of CVDgrown diamond films on different substrates. Therefore, it is critical to investigate the parameters that control diamond films' adhesion on substrates' surfaces and develop processes to produce significant improvements on diamond films adhesion on many surfaces for different diamond-coated products [180, 181]. A process that can potentially improve diamond films' adhesion on substrates is the introduction of a template layer on the surface of the substrate, which can induce an enhanced chemical attachment of the diamond film [182, 183]. The potential mechanisms associated to the template layer for improving diamond films' adhesion include: (1) enhancement of density of diamond nucleation areas on the substrate surface; (2) minimization of thermal and interfacial stresses; (3) improvement of the surface chemistry of the template layer to induce stronger chemical bonding of the diamond film.

Polishing and planarization of PCD films

The surface roughness and film thickness poor uniformity may induce limitations in different applications of CVD diamond films, induced by different performance parameters, namely: a) insufficient thermal contact for thermal management at the diamond/surface interface, b) light reflection at surface of diamond-coated optical windows, undesirably reducing light transmission, c) lack of reproducibility for electrical contact of metal electrodes on the surface for diamond based electronic active devices, and other effects. MCD films' high surface roughness induce limitations on the smallest size achievable for diamond films-based MEMS/NEMS devices. The limitations on applications of diamond films for fabrication of many MEMS/NEMS devices, based on surface roughness, can be strongly reduced or eliminated by polishing the diamond surface to achieve a desirable surface roughness for a specific application. Many techniques have been developed and are being used to polish diamond films grown by the CVD process, namely: (1) mechanical polishing [184], (2) polishing via complementary thermo-chemical processes [185], (3) mechanical polishing assisted by complementary chemical process [186], (4) polishing by laser beam swapping over the diamond film surface at grazing incidence [187], (5) polishing by ion beam bombardment of the diamond surface at grazing incidence [188], (6) reactive ion etching (RIE) [189], and (7) liquid jet abrasion-induced polishing with solid particle impact [190, 191]. The characteristics of various diamonds polishing techniques are summarized in [192] where the details of each polishing method are illustrated.

Polished MCD film surfaces may not be flat at the nanoscale level. In addition, secondary effects, produced during polishing, such as formation of submicron-cavities between diamond crystals in PCD films' grain boundaries, generated during the synthesis of the PCD films, are opened as surface pits during the polishing process. To achieve a nanoscale level smooth surface, these micro-cavities must either be reduced or eliminated during growth (ideal solution) or filled following the polishing process. Organic monomer photoresist, spin-on-glass (SOG), or polyimide may be used as a filler material to fill the cavities [192]. These filler materials are polymerized when heated at temperatures in the range 80–425 °C, yielding a smooth surface. A substantially better approach to produce PCD films with smooth surfaces, without requiring the lengthy and costly polishing procedures described above, is provided by the growth of UNCD films, which exhibit surface roughness of 3–5 nm, similar to the grain size of these films (see detailed description in Sect. 10 of this review and a detailed review of the UNCD film technology published in ref. [4].

Processes for miniaturization of PCD filmsbased MEMS structures

In relation to how small diamond-based MEMS/ NEMS structures can be fabricated, PCD films-based nano-resonators were fabricated as shown in Fig. 18a, using e-beam lithography [193]. The fabricated nanoresonator structures included cantilever structures patterned to have widths as narrow as 200 nm (Fig. 18a), and torsional paddle resonators with a support beam width of 0.5 µm (Fig. 18b). The performance of these structures shows resonant frequencies and Q values in the range of 23-805 kHz and 9500-103,600 respectively. Recently, the world's first SCD nanoelectromechanical switch was reported by Japanese researchers as shown in Fig. 19. The developed devices can be applied as microwave switch for next-generation wireless communications and logic circuit under harsh environments. The NEMS devices described above establish the infrastructure for producing diamond based NEMS with novel functions, opening the way for the development of various chemical, physical, and mechanical sensors. In relation to producing SCD film-based

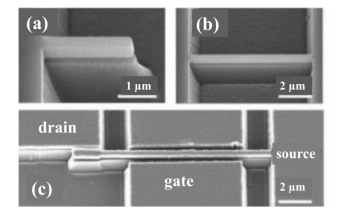


Figure 19 SEM images of suspended structures of SCD film based NEMS cantilevers: **a** cantilever, (**b**) bridge, and **c** 3-terminal NEMS switch. Air gap structure has been formed in the substrate side [274]. Notice that the cantilever structures are not lifted from the substrate.

NEMS structures, as shown in Fig. 19, an important process that needs to be developed, when growing SCD films on SCD substrates, as show in Fig. 19, is etching of the SCD substrate, to release the SCD filmbased cantilever.

Diamond-based MEMS sensors, actuators and other devices

Diamond Film-based SAW MEMS devices

Surface acoustic wave (SAW) devices, representing a group of sensors based on resonant frequency's detection as the sensing mechanism, have

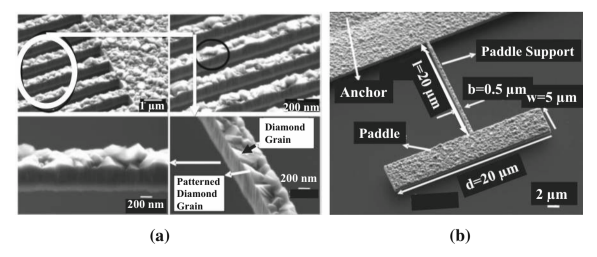


Figure 18 SEM image a shows smooth sidewalls, but rough top surface MCD film-based nanocantilevers fabricated by e-beam lithography and b MCD film -based single torsional resonator [193].



demonstrated good efficiency in detecting bio-species such as DNA and proteins. In this sense, the high acoustic phase velocity of synthetic diamond makes it very attractive for the achievement of high frequency (GHz range) SAW devices when combined with suitable electric materials. Among available piezoelectric materials, Aluminum Nitride (AlN) [194–197] and Zinc Oxide (ZnO) [198-200] has been generally used for high frequency SAW diamond devices. As a piezoelectric material, AlN is preferred owing to its high acoustic phase velocity and its fairly large piezoelectric coupling coefficient along *c*-axis [197]. Among diamond materials, NCD is preferred compared to MCD because its nanometric grain size could help reduce the important propagation losses usually encountered in MCD [201]. There are two possible configurations of layered structure of a SAW device depending on the placement of (IDT) inter-

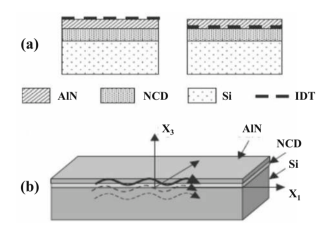


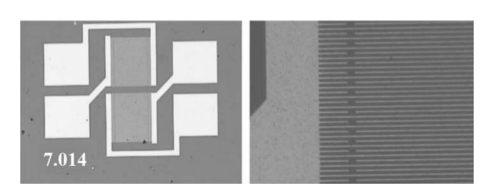
Figure 20 a Cross-sections schematics of two possible IDT configurations for an AlN/NCD/Si layered structure. b Schematic diagram of the AlN/NCD/Si layered structure together with the system of coordinates considered in the propagation model [196].

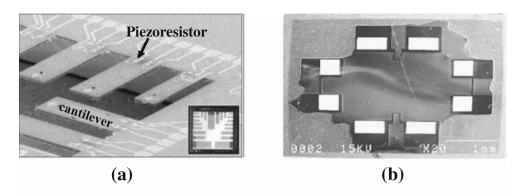
digital transducers as shown in Fig. 20. A fabricated diamond-based SAW device is shown in Fig. 21. A SAW device with such layered structure was reported with operating frequency of 5 GHz and acoustic phase velocity of 9.2 km s⁻¹.

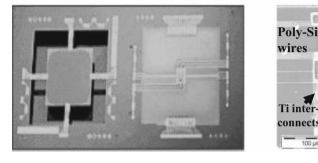
Diamond film-based Piezo-resistive MEMS sensors

Piezo-resistive-based Si technology is widely used for pressure, force and acceleration-based MEMS sensors. A widely used device based on piezo-resistivity effect is the piezo-resistive acceleration MEMS sensor that deploy airbags in cars during an impact related accident. Commercially available sensors cannot operate at high temperatures because of pn-insulation of the piezo-resistors. The first report of piezoresistivity in SCD and PCD films-based MEMS sensors, by researchers at Michigan State University in 1992 [202], generated substantial interest in the utilization of p-type PCD films grown by chemical vapor deposition (CVD), as a material for sensor applications, owing to its unique physical and chemical properties and potential operation at high temperatures. A number of studies focused on the piezo-resistive effect of PCD films-based sensors [203–216], such as the gauge factor as a function of operating temperature. A MEMS cantilever or diaphragm made of either Si, stainless steel or undoped PCD film, is usually used as a structural material. The gauge factor is usually determined by measuring the change of piezo resistor by pressing down the structural material. Studies demonstrated that the gauge factor of PCD films increases from 5 to 100, at room temperature, with the increase of resistivity, grain size and operation temperature. The reported gauge factors of PCD films in the literature were already shown in Table 5

Figure 21 Example of SAW device fabricated with an AlN/ NCD layered structure, via direct writing with e-beam lithography [197].







(c)

Ti interconnects

Poly-C tip sensor

Figure 22 a Diamond-based MEMS cantilever test structure to characterize the gauge factor [213], **b** a pressure sensor using p-type PCD film-based piezo-resistor [222], **c** PCD film-based

acceleration sensor (left) and pressure sensor (right) [213], **d** position sensor made of p-type PCD film-based cochlear implant probe [71].

Figure 22a shows a cantilever made of undoped PCD film, used as the structural material to characterize the gauge factor of p-type PCD film-based MEMS piezo-resistor. Among the applications of PCD film-based piezo-resistive sensors, pressure sensors have been the most investigated [217–221].

Figure 22b and c show pressure sensors and an accelerometer from the work described in references [214, 221]. Recently, a position sensor using PCD film-based MEMS structure was developed for insertion in a cochlear implant probe [71] as shown in Figure 22d. RF-MEMS resonators with piezo-resistive detection are also being developed [222, 223]. The piezo-resistive detection, related to device scaling, without decreasing the detection sensitivity, can help adjust the output impedance by: (a) adjusting the piezo-resistor dimensions and (b) increasing the mobility of holes by using intra-grain piezo-resistor. It has been observed also that DLC films also exhibit the piezo-resistive effect and pressure sensor and force sensor made of DLC films have been reported [224–228]. Since the Young's modulus of diamond is much higher than for Si and metals, diamond-based piezo-resistive devices can operate at substantially higher frequencies, as discussed in Sect. 6.1

PCD films-based MEMS gas sensors

Diamond films can also behave as chemical sensors measuring hydrogen, oxygen and many other chemicals' concentration. For example, diamondbased chemical gas sensors, using Pd/undoped PCD film/highly-doped PCD metal-insulator-semiconductor (MIS) diode structure were made and the hydrogen sensing characteristics investigated as a function of hydrogen partial pressure and temperature [229–232]. The physical operation of a hydrogen sensor involves effects as described briefly next: (a) H_2 molecules from the environment adsorb on the surface of the Pd electrode; (b) H_2 molecules break apart when interacting with the surface of the metal electrode, and diffuses through the palladium (Pd) metal electrode layer to the metal / insulator interface; (c) H atoms adsorb at the undoped PCD film-



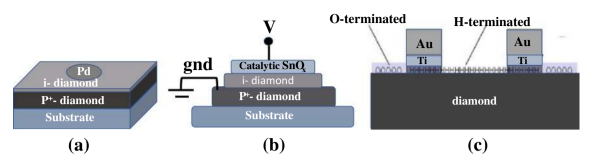


Figure 23 a hydrogen sensor using Pd/undoped PCD/highly doped PCD metal-insulator-semiconductor (MIS) diode structure [233], b oxygen sensor using Pt/SnO_x/undoped PCD/highly doped

PCD MIS diode structure [235], c gas sensor using hydrogenterminated diamond [236].

metal interface, creatin a Pd metal electrode dipole layer; (d) the dipole layer induces a transformation of the work function difference between the Pd metal electrode layer and the undoped PCD film; (e) the relation between the current and the applied voltage to the MIS diode changes; and (f) changes in the current-voltage curve may be induced by the H₂ gas partial pressure in the environment in contact with the device [233]. The typical structure of a H- sensor is shown in Fig. 23a [230]. A diamond-based chemical gas sensor using Pt/SnO_x/undoped PCD films/ highly-doped PCD film MIS diode structure, for oxygen sensing, was also reported [234, 235]. Research provided evidence indicating that atomic O atoms can diffuse into the bulk of PCD films, from the surface of the tin oxide, leading to a decrease in the number of O atoms vacancies in tin oxide, resulting in a decrease in the electrical conductivity of tin oxide. This decrease in the electrical conductivity of tin oxide modifies the voltage distribution across the MIS structure, which, in turn, decreases the diode current. This layered structure can also detect CO gas. A characteristic structure of an O gas sensor is depicted in Fig. 23b [235]. The Pd-gated diamondbased MIS sensor is also sensitive to benzene (C_6H_6) and toluene (C_7H_8) gases [233]. The information presented above indicates that benzene and toluene are adsorbed as some forms of hydrocarbon radicals. Besides the use of MIS structure for gas sensing, hydrogenated surface layer of diamond could also be used as shown in Fig. 23c, due to the surface transfer doping mechanism [236]. The steady-state gas response is only limited to analyte gases that can undergo electrolytic dissociation in the surface liquid electrolyte layer and can be enhanced by a small number of O-termination sites.

PCD Films-based UV, X-ray and particle detectors

Diamond is also an excellent material for UV detection [237-241], x-ray detection [242-244] and particle detection, including neutron detection [245, 246], alpha particle detection [247], proton detection [248] and single-ion detection [249-251]. The 5.5 eV band gap of diamond induces an extremely low dark current and specific absorption of $\lambda < 225$ nm wavelight. In addition, diamond exhibits length outstanding electronic properties and strong resistance to radiation. Many detectors of the type mentioned above are made from intrinsic diamond instead of p-type doped diamond. The two main designs for diamond-based sensors feature top and bottom electrode layers encapsulating the diamond layer, or interdigitated electrode-diamond layers, as shown schematically in Fig. 24 [252]. The mechanism of UV, X-ray or particle detection is as described in the following text. A charged particle, or a photon with energy above the bandgap, passes through the diamond and ionizes C atoms, leaving a trail of electron-hole pairs, which are separated by the electric field applied between the electrodes. Some electrons are trapped within the diamond PCD film at defects and grain boundaries and may contribute to space charge build up inducing polarization of the crystal. Other electrons are inserted in the electrodes, contributing to the photoconductive current in the detector. Current CVD methods produce PCD films with large grain boundaries' networks and defects in the crystal structure of the grains and in grain boundaries, affecting the PCD films photoconductivity. Han and Wagner [253] showed that grain boundaries of PCD films degrade the detection properties by a factor of two to three when

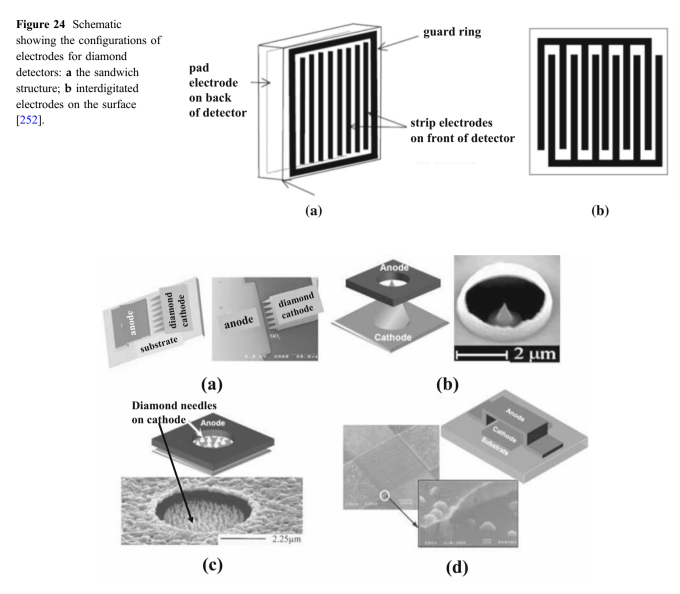


Figure 25 Overview of field emission devices based on PCD films: (a) Lateral emitter structure [258], (b) vertical emitter structure with PCD micro-tips fabricated by mold transferring technique [262], (c) vertical emitter with diamond micro-tips

fabricated by selective growth methods [263] and (d) vertical emitter that allows a small distance between the cathode and the anode with a precision limited by lithographic resolution [266].

interdigitated electrodes are used, since carriers are more easily trapped than drifting across grain boundaries.

PCD field emission devices

There has been an increasing interest in developing PCD film-based field-emission (FE) devices due to their huge application potential and lower cost. Diamond is used for field emission because key properties: (1) low operating voltage due to its negative or low electron affinity, (2) constant emission current

due to the high chemical stability, and (3) the stability of structure due to its superior hardness. In order to permit (a) lower operating voltage, (b) high accuracy of anode to emitter spacing, and (c) well defined emitter area, the recent research focus has been on devices with a built-in anode. PCD film-based fieldemission devices include lateral electron field emitters [254–261] and vertical gated electron field emitters [262–266]. Among these structures, the cathode is made of diamond with or without micro-tips. The anode is either metal or Si.

An overview of PCD films- based emitter structures, including the new structures used in the current work reported in this paper, is shown in Fig. 25, including concept diagrams. A typical reported lateral emitter structure is shown in Fig. 25a, which is subject to problems related to (i) creation of very small spacing between the anode and the cathode, and (ii) the difficulty of achieving sharp tips. Both of these factors may result in a high operating voltage. The reported vertical gated FE structures are of two types. One type contains diamond micro-tips fabricated by either mold transferring techniques [254, 262] or selective growth methods [263, 264 with marginal emission characteristics, as shown in Fig. 25b, c, respectively. Compared to the lateral electron emitter, the vertical gated electron field emitter can have very sharp tips to achieve high field enhancement factors, whereas the distance between the tip and the circular shaped anode may still be large. Figure 25d shows another structure [265, 266] which allows a small distance between the cathode and the anode with a precision limited by lithographic resolution.

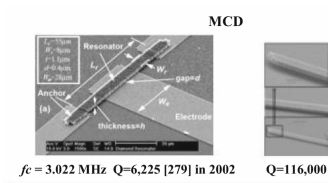
Diamond films-based RF-MEMS

Radio frequency micro electromechanical systems (RF-MEMS) can yield on-chip micromechanical resonators with ultrahigh quality factors over 10,000 at GHz frequencies in both vacuum and air, making them excellent candidates for broadband wireless communications [267]. In addition, RF-MEMS resonators can be excellent mass and chemical sensors, specifically by controlling the chemical surface termination. So far, polycrystalline silicon has been the preferred material for RF-MEMS resonators, having been demonstrated with measured quality factor Q's above 8400 at a frequency of 50.35 MHz [268] for freefree beam design, Q's of 2650 at a frequency of 1.156 GHz [269], for radial disk design, Q's of 2800 at a frequency of 1.52 GHz, for extensional wine-glass design [270, 271]. The impressive outcome of the research described above, can be greatly enhanced if Q's > 10,000 were achievable at the same GHz frequencies and in the same tiny sizes. In order to further extend device operating frequencies, the use of alternative structural materials, with higher acoustic velocities, given by the formula $\sqrt{E/\rho}$ (E:Young's modulus and ρ : material density), like diamond and

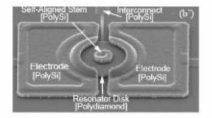
SiC [272] 273, over Si, has been explored. Among the currently available set of thin-film materials, diamond offers the largest acoustic velocity on the order of 18,076 m/s [271]. This is to be compared with the 8,024 m/s of single crystal silicon [272] and 11,500 m/s of silicon carbide (SiC) [273], which are 2.25X and 1.57X smaller, respectively. Given that resonance frequency is generally proportional to acoustic velocity, diamond provides the largest boost towards even higher MEMS-based resonator frequencies. In addition, the electrical and mechanical properties of polycrystalline Si begin to rapidly degrade at temperatures above 350 °C, making it increasingly unsuitable for devices required for high temperature operation.

Fabrication and testing of diamond-based RF-MEMS resonators have been reported in the literature. They were made of SCD [274], MCD [275-278], NCD [29, 279-283], UNCD [4, 284] or DLC [285, 286]. Figure 26 shows pictures of some reported diamondbased RF-MEMS resonators and their corresponding quality factor Q's at resonant frequencies f_0 's. More quantitative performance results are given in Table 9. The dissipation (affecting *Q*) is principally induced by the relaxation of substantial number of defects in the bulk or surface of the PCD film. Sepúlveda et al. [277] reported a quality factor Q of 4000–100,000 for PCD films (grain size \sim 300 nm)-based cantilevers, which includes some values higher than the Q's observed for UNCD cantilevers with comparable dimensions [286]. They proposed the concept that as the percentage of the nucleation layer containing fine grained diamond is increased, the quality factor Q of the resonators is reduced. This result correlates with the hypothesis that higher dissipation observed in UNCD and DLC films is principally due to the presence of the larger number of grain boundaries and defects. Low quality factor Q of NCD in Table 9 may arise from the fact that as the resonator dimensions become smaller, dissipation due to surface defects may increase. This result is supported by research performed by Imboden et al. [280]. From many experiments, it seems that SCD is potentially the best material top achieve high Q. However, the problem is that growth of SCD films on large area substrates for low coast industrial fabrication of SCDbased sensors is not currently suitable as for PCD films-based devices. Thus, PCD film-based RF-MEMS resonator appears to be the best candidate to achieve higher quality factor Q beyond Si for now.

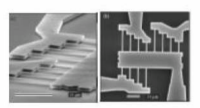
Figure 26 Typical RF-MEMS resonators made of SCD, MCD, NCD, UNCD and DLC films.



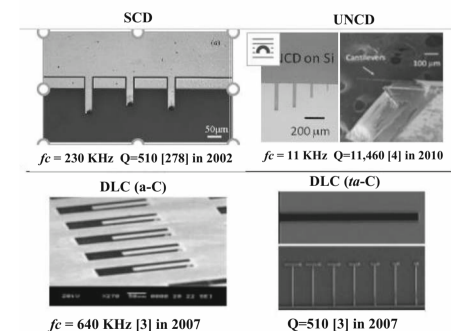




fc = 1.51 GHz Q=11,555 [279] in 2004



fc = 17.66 MHz Q=600-2,400 in 2007



fc = 640 KHz [3] in 2007

A classical electrostatically driven cantilever beam structure switch is shown in Fig. 27 [287, 288]. The main part of the device is a freestanding diamond cantilever that can be deflected electrostatically by

RF-MEMS switches exhibit properties with great potential to enable a new generation of RF communications devices, because of high isolation and low power consumption enabled by the device structure.

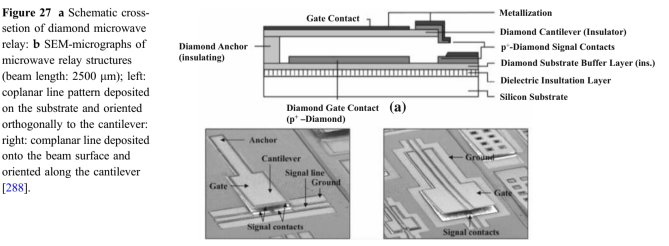
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Resonator design	Quality factor Q	Measured frequency f_0	Actuation method	Mater
Cantilever [275]	510	∼ 230 kHz	Piezoelectric	SCD
Doubly clamped beam [276]	6225	3.022 MHz	Electrostatic	MCD
Comb drive [278]	36,460	27.352 kHz	Electrostatic	MCD
Cantilever [277]	15,260	384.9 kHz	Piezoelectric	MCD
Cantilever [278]	116,000	8–50 kHz	Piezoelectric	MCD
Doubly clamped paddle [29]	2400-3500	6–30 MHz	Piezoelectric	NCD
Mesh membrane [29]	3000	8–20 MHz	Piezoelectric	NCD
Disk resonator[279]	11,555	1.51 GHz	Electrostatic	NCD
Doubly clamped beam [282]	~ 3000	14–157 MHz	Magnetomotive	NCD
Tuning fork [283]	8000	37.0 MHz	Laser	NCD
Square paddle (pillared) [283]	1500	14.77 MHz	Laser	NCD
Square paddle (freely suspended) [283]	3000	13.85 MHz	Laser	NCD
Ring resonator [283]	5000	40.18 MHz	Laser	NCD
Antenna structure [284]	23,200	630.6 MHz	Magnetomotive	NCD
Harp structure [281]	600-2400	17–66 MHz	Magnetomotive	NCD
Cantilever [4]	11,460	11 kHz	Piezoelectric	UNCD
Cantilever [285]	5000-16,000	12–35 kHz	Piezoelectric	UNCD
Cantilever [286]		108 MHz	Piezoelectric	DLC (a-C)
Cantilever [280]	3500	KHz	Piezoelectric	DLC (ta-C)

Table 9 The reported room temperature quality factor Q of flexural resonators

applying a voltage between the cantilever gate contact and the substrate gate contact. In the deflected sate, the cantilever is bent towards the substrate and current flow across the cantilever signal contact and the substrate signal contact. In the off state the signal contacts are separated by an air gap. All essential parts of the switch are made of diamond except the Si substrate. Thus, heat generated caused by the power loss in the signal contacts can be effectively dissipated over the cantilever and the substrate. Also, no sticking caused by the melting of metal contacts or

formation of insulating oxide layers on the signal contacts can occur. An on-state attenuation of ~ 3 dB and an off-state attenuation of - 23 dB at 10 GHz were reported [288]. The results indicate that, for operation in air, a diamond microswitch exhibits approximately eight times higher maximum frequency of operation than a Si microswitch of identical geometry [289]. The disadvantages of it are high driving voltages and low contact force due to the large Young's modulus of diamond. Alternatively, the bi-metal effect has been employed in an



[288].

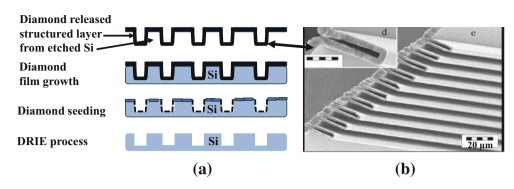


Figure 28 a Cross-cut view of 4 µm tunnels, b fabrication process of microfluidic channels [3].

electrothermal driving concept. In this concept, the cantilever is bent thermally induced stress. The actuation voltage can be as low as 1.3 V, which is significantly lower than the actuation voltages of electrostatic switches, which are in the order of several 10 s V. However, thermal power is applied to close the switch and is also needed to keep it closed, which can produce a permanent power loss. To avoid this disadvantage a bi-stable configuration had also been developed based on a prestressed double-anchored beam [290].

Diamond-based BioMEMS microfluidic channels

Heat dissipation is currently a critical challenge to be overcome by the microelectronics industry. Miniscale coolers involving cooling fluid passing through micron-size channels provide and advanced technology to extract thermal energy from microelectronics circuits such as CPUs. Microfluidic channels are also key components of lab-on-a-chip systems enabling complex chemical/biochemical processes as well as analytical tasks. Ramesham, et al. used anisotropic chemical etching of silicon and selective growth of diamond to form PCD films- based microfluidic channels of 17-µm wide [291]. This processing resulted in channels with diamond forming the top half of the structure and silicon the bottom half. Such channels may not be useful in many microfluidic applications due to the relative chemical reactivity and thermal instability of silicon compared to diamond. Closed surface channels made of PCD films, with a sufficient large cross-section, were reported by Müller, et al. using electroplated copper as mold and sacrificial layer [292]. The copper layer was finally etched by sulfuric acid. A micro membrane pump system was fabricated and characterized based on this process. Guillaudeu, et al. developed free-standing all-diamond fluidic channels using Si as a mold as shown in Fig. 28a [293]. This process is simpler but only narrow width can be designed to close the trench in a proper time by additional PCD film growth as shown in Fig. 28b. The fabrication process, developed by Guillaudeu et al. [293], was modified to integrate an active diamond heating element within the cavity [294]. This process enables the fabrication of an "all-diamond" bubble-jet-based device, for which the fluid is in contact only with diamond as shown in Fig. 29. A B-doped diamond underneath the channel structures was used as a heating element for the bubble-jet element.

MEMS neural probes

Many groups are performing R&D to develop industrial-type processes for fabrication of neural probes. In relation to these efforts, the material needs to exhibit the best possible biocompatibility. In this sense, Si is not an appropriate material, since exhibits poor flexibility, solubility in water, and can induce undesirable glial responses [294]. Several groups have demonstrated an excellent potential solution to this problem, through coating silicon with diamond coatings, which exhibits unique mechanical, tribological, chemical properties and high biocompatibility [4, 295, 296], and as discussed in Sect. 10. The optical transparency of diamond, used as a substrate, is important for in vitro experiment because it allows the electrode's position on the probe to be easily located under a microscope. Recently, B-doped PCD films-based sensors and electrodes were integrated in Si cochlear microprobes for the first time [297]. B-doped PCD films-based electrodes have also been

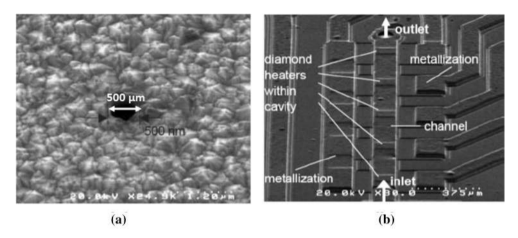


Figure 29 a SEM image of a nanoscale hole (500 nm diameter) etched in a diamond membrane, b peristaltic pump of 5 diamond heaters in series within a channel. [294].

shown to be effective for the detection of neurotransmitters [298]. Chan et al. [299–301] developed a process for fabricating neural probes using undoped PCD films as the structural material and B-doped PCD films as the electrode material. NCD films are being implemented in interconnects and electrical insulation layers, in probes used to measure electrical neural activity. In addition, the comparatively wide potential window (reported values range from 1.4 to 4 V) in an aqueous environment [302, 303], low double layer capacitance (ranging from 5 to 40 μ F/ cm²) [303], chemical inertness and stability, resistance to fouling of B-doped diamond makes such PCD film-based probe an excellent material for electrochemically detecting neurotransmitters in vitro as shown in Fig. 30 [304, 305].

PCD film encapsulation of Si-based MEMS devices

Although MEMS packaging may benefit from mature packaging techniques from the semiconductor IC industry, MEMS packaging is still complicated due to the diversity of applications. PCD films have emerged as a novel material for MEMS [304–307]. The potential of PCD as a thin film packaging material was explored in view of its excellent mechanical strength, electrical properties, chemical stability and thermal management as a heat sink. A PCD thin film packaging process was developed to encapsulate cantilever resonators [307] integrated with highly doped PCD film interconnects [307], as shown in Fig. 31. The efficiency of using a PCD film

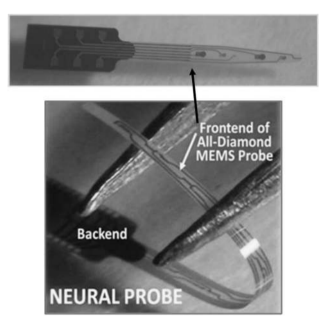


Figure 30 PCD-based probe used for electrochemically detecting neurotransmitters [299].

encapsulation process was evaluated by measuring the resonator frequency and quality factor before and after the packaging process, which indicated that PCD film-based packaging may be integrated into conventional MEMS fabrication without affecting the devices' performance yields. The hermeticity for the fluidic performance was also tested by an HF soak test, which revealed that PCD films encapsulated cantilevers can perform in harsh environments, withstanding chemical and mechanical attack and high temperatures. A technology using an all-diamond package for wireless-integrated MEMS with

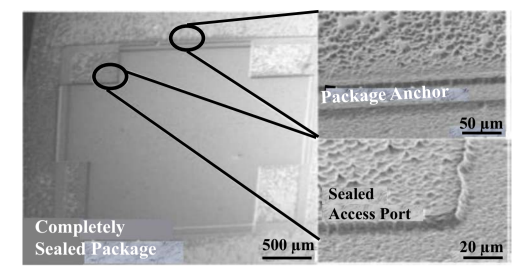


Figure 31 Completed sealed PCD film packaging [307].

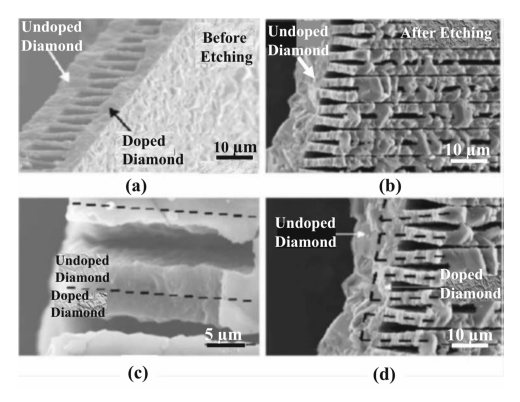


Figure 32 SEM image of PCD films encapsulated panel with built-in interconnects: a PCD film encapsulated panel before dry etching, b PCD film encapsulated panel after dry etching, c top

view of PCD film encapsulated panel, and **d** side view of PCD film encapsulated panel [307].



B-doped PCD films, as built-in interconnects was also demonstrated [307] (Fig. 32), showing that PCD films based MEMS devices may exhibit an outstanding level of heat dissipation.

Ultrananocrystalline diamond (UNCD) film technology for environmental and human body implantable MEMS devices

Brief Introduction to UNCD film technology

UNCD film growth via microwave plasma chemical vapor deposition (MPCVD) using conventional diamond nanoparticle-based seeding of substrate surfaces

The discovery of a new process to grow polycrystalline diamond films with grain sizes < 10 nm yielded the breakthrough represented by the ultrananocrystalline diamond (UNCD) film technology. The field of UNCD film based technology started with the discovery by Gruen/Krauss/Auciello that replacing the H_2 gas in the H_2/CH_4 gas mixture, used previously to grow from SCD to MCD to NCD films [4, 27, 34, 35], by Ar gas (the less expensive inert gas in the market today). to flow an Ar (vol. 99%)/CH₄ (vol. 1%) gas mixture in the MPCVD system, resulted in the growth of polycrystalline diamond films with grain size in the range of 3–5 nm size, the smallest grain size of any polycrystalline diamond film today

(see reviews [4] and patent [33]). Briefly, the original R&D, using the MPCVD film growth technique, demonstrated that diamond films grown by MPCVD, using an Ar (99%) / CH_4 (1%) gas mixture flown into an air evacuated chamber of an MPCVD system, and coupling microwave power, produced a green plasma involving C_2 (C=C) dimers, with no H atoms bonded, thus enabling an extremely $low(\sim 6$ kcal/mol) activation energy for C atoms sp^3 (diamond) bond formation on substrate surfaces. acting as the main UNCD nucleation species to induce nucleation and contribute to film growth, jointly with CHx (x = 1, 2, 3) species created also in the plasma. In this sense, recent modeling [308] indicated that although the C_2 content in the plasma is high, the density near the substrate surface may be lower, and other hydrocarbon radicals (e.g., CH₃, C₂H₂) may also contribute to the growth of UNCD films. However, the model [308] is related to the grow of diamond films produced by MPCVD, using a mixture of Ar/CH₄/H₂ gases, which does not produce the unique UNCD structure (with 3-5 nm grains) grown by MPCVD with the Ar/CH_4 chemistry [4, 33]. In addition, the model [308] does not explain the low temperature growth of UNCD films as demonstrated for UNCD film growth at 350-400 °C [309, 310]. In any case, diamond films with 3–5 nm grain sizes (see Figs. 33a,b), which induced the coining of the name UNCD, with surface roughness of \sim 3–5 nm rms (see Fig. 33c), lowest coefficient of friction (compared with MCD, NCD, and DLC (see Fig. 33c), practically

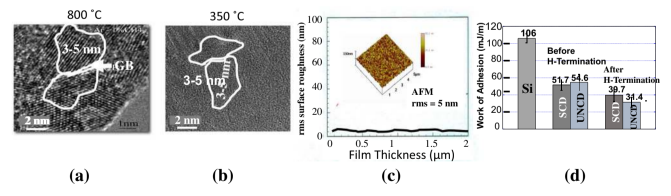


Figure 33 a HRTEM image of a UNCD film grown at 800 °C, showing the 3–5 nm grain size; b HRTEM image of a UNCD film grown at ~ 400 °C (lowest temperature demonstrated for growing polycrystalline diamond films), showing the 3–5 nm grain size; (c) rms surface roughness of UNCD films vs film thickness, showing that the surface roughness of UNCD films is practically independent of film thickness and the smoothest

compared with any other as grown PCD film; (d) Work of adhesion of Si, single crystal diamond (SCD) and UNCD films surfaces, showing that UNCD films exhibit, overall, the best low work of adhesion to enable MEMS/NEMS devices with components in surface contact, without performance degradation due to surface stiction.

no stiction in contact with other materials (see Fig. 33d), excellent dielectric properties, and no mechanical wear [4, 33], enable a new generation of superior MEMS/NEMS devices, as described in the following sections.

The MPCVD-based UNCD film growth process described above is based on prior conventional process of seeding the surface of the substrate with micro- or nano-diamond particles, to induce the nucleation and growth of the UNCD films. The most used seeding process today, involves immersing the substrate in a container, with a solution of micro- or nano-size diamond particles in methanol, in an ultrasound wave generating system, such that the sound waves shake the diamond particles, embedding them on the substrate surface as "seeds" to induce the nucleation and subsequent growth of diamond films (see review [4]).

UNCD film growth using MPCVD bias enhanced nucleation-bias enhanced growth (BEN-BEG) process

A new approach to grow UNCD or any other PCD films, using the MPCVD process, eliminating the extra costly step of the chemical seeding process, involves biasing electrically conductive or semiconductive substrates, like Si, with a negative voltage with respect to the plasma, to accelerate C^+ and CHx^+ (*x* = 1, 2, 3) ions by 100–300 eV, which upon impacting on the substrate induce Carbide layer formation, nucleating the diamond films, and subsequently contributing to the growth process. Early research on exploring the concept of BEN-BEG of UNCD films was done first when exploring growing UNCD films with CH₄/N₂/very small H₂ gas mixtures in an MPCVD process [311], to produce N-UNCD films with N atoms in grain boundaries to develop electron field emission devices [311]. This research provided first experimental evidence supporting earlier theoretical work [312] proposing a mechanism for BEN of diamond films Subsequently, two groups demonstrated BEN processes to grow UNCD films using the patented [33] CH₄/Ar gas mixture in the MPCVD process. One group demonstrated that BEN of UNCD films, followed by growth without bias on Si substrates [313], exhibit stronger adhesion to the Si substrate's surface (a very important effect for UNCD film-based MEMS/NEMS devices) than for UNCD films grown on chemically seeded Si surfaces [4, 309, 310]. The stronger diamond film adhesion to substrates, produced by the BEN-BEG process, is induced by the relatively high kinetic energy of C^+ and CHx^+ ions extracted from the plasma and accelerated to the Si substrate surface, which easily form covalent bonding, Si–C, and bond strongly to both the Si and subsequently to the growing UNCD film.

Auciello's group performed systematic R&D to develop a low-pressure BEN-BEG process to grow UNCD films [314]. using the MPCVD-based BEN-BEG technique, involving key steps, namely: (1) etching of the natural SiO₂ layer on the surface of a Si (100) substrate, for 10 min, in a pure H-based plasma, with a substrate biased at -350 V; (2) in situ BEN-BEG, using a H_2 (93%)/CH₄ (7%) gas mixture, to involve H⁰ atoms and H⁺ ions in the plasma for etching the impurity graphitic phase that in many cases growth concurrently with the diamond phase when growing NCD films [315-317]. The grow of UNCD films by BEN-BEG involved a plasma produced by 2.2 kW microwave power at low 25 mbar pressure and applying -350 V bias on a substrate heated to 800 °C in a 2.45 GHz 6 in. IPLAS CYRANNUS MPCVD system [314]. As the UNCD film nucleates, the electron emitted current from the substrate increase to a stable value (Fig. 34a), while the surface roughness of the films decreases from the low NCD (~ 10 nm), while grains are nucleating (Fig. 34b), to the UNCD surface roughness of ~ 6 nm (Fig. 34a), when the UNCD film reach high density (Fig. 34c), with the surface roughness measured by high resolution AFM (Fig. 34d). HRTEM showed that the ion bombardment, during the BEN process, induces formation of pyramidal Si structures (Fig. 34e), produced by preferential ion beam sputtering along the Si {100) crystallographic direction. Subsequently C atoms bond chemically with Si atoms on the tip of the pyramids nucleating initial interface SiC layer, on which the UNCD film grows, while some initial graphite interface layer develops in between the Si pyramids (Fig. 34f) and related electron diffraction patterns), subsequently also enabling UNCD film growth (Fig. 34f). The BEN-BEG process described above yielded UNCD films with low stress, smooth surfaces (\sim 4–6 nm), high growth rates $(\sim 1 \,\mu\text{m/h})$ (see Fig. 34c and uniform grain size (3-7 nm) throughout the whole film area on 100 mm diameter Si wafers.



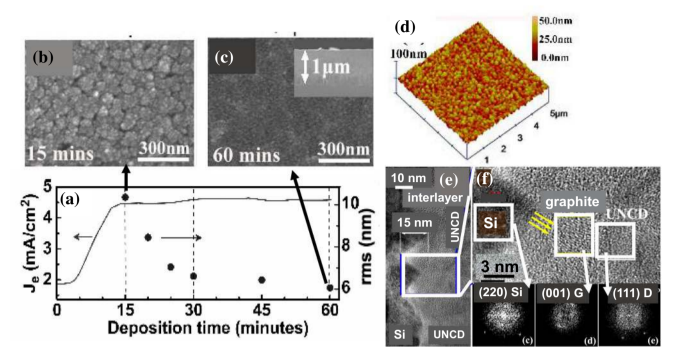


Figure 34 a Plasma electron density current versus UNCD film growth time during BEN + BEN-BEG process; **b** SEM image of initial UNCD grain nucleation on Si substrate; **c** SEM image of UNCD film after 60 min growth time, when UNCD film reaches 1 μ m thickness with dense structure with ~ 6 nm grain size; **d** (mixi o) high resolution AFM image obtained when measuring

UNCD film growth using hot filament chemical vapor deposition (HFCVD) Process

UNCD film growth via HFCVD using conventional diamond nanoparticle-based Seeding of Substrate Sur*faces* Recently, Auciello's group performed systematic research that demonstrated that UNCD films can also be grown on large area substrate with excellent density, nanoscale grains (\sim 3–5 nm) and thickness uniformity across 100 mm diameter Si wafers using the HFCVD process [318]. The HFCVD process involves setting up an array of nine parallel W filaments held on a Molybdenum (Mo) frame (Fig. 35a,b), positioned above the substrate holder at a distance of \sim 3 cm (Fig. 35a). The substrates are positioned on a Mo disk, which is rotated during film growth to induce film thickness uniformity on up to 100 mm diameter substrates. The filaments are heated to ~ 2200 °C, via passing an AC current during the film growth. Several gas ratios were investigated for diamond film growth, using the conventional nanocrystalline diamond seeding process described for the MPCVD process. Table 10

UNCD surface roughness ~ 6 nm rms; **e** and **f** cross-section HRTEM image of Si substrate showing Si tips, produced by plasma Ar + , C⁺, CHx⁺ ions bombardment, inducing nucleation of SiC (on top of Si tips) and graphite in between Si tips, plus UNCD initial layer, all confirmed by electron beam diffraction, shown at the bottom of **f**.

shows the gas mixtures flown to grow MCD, NCD and UNCD films. The H_2 (10 sccm)/CH₄ (2 sccm)/Ar (90 sccm) gas mixture produced UNCD films with nanostructure identical to those shown in Fig. 33a,b. The systematic research on the HFCVD process [318] provided substantial improvement on understanding the HFCVD process to grow UNCD films, previously investigated by other groups, which was limited in extend and deepness [319, 320].

UNCD film growth via HFCVD bias enhanced nucleation-bias enhanced growth (BEN-BEG) Process Growth of MCD to UNCD films using the aggressive "seeding" process in an ultrasonic system can damage most 3-D micro/nanostructured surfaces, such as micro/nano-rods of 3-D nanoscale Si or other materials being investigated to produce diamond-based MEMS nanostructured components. Therefore, the development of a BEN-BEG process using HFCVD is, as the MPCVD BEN-BEG process, a key diamond film growth process for a new generation of diamond and particularly UNCD film-based MEMS/NEMS devices.

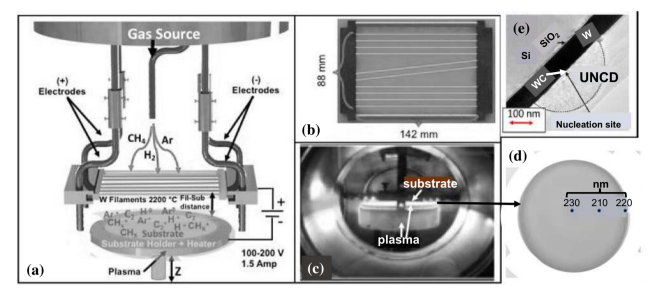


Figure 35 a Schematic of the W filament array/substrate used for the HFCVD BEN-BEG process described in Ref. [324]; **b** Schematic of the top view of the filament array, featuring two central filaments at an angle to equilibrate the density of C and CHx species production, from cracking of CH₄ molecules on the hot filament's surface, with respect to similar species produced from the parallel filaments towards the substrate edge, to produce uniform UNCD films all across the substrate surface; **c** HFCVD in operation growing a UNCD film by BEN-BEG; **d** Optical image

Table 10 Effect of gasmixture flows on grain size ofPCD films grown by HFCVD

of a UNCD film grown on a W coated (70 nm thick) high electrically conductivity Si substrate, showing excellent film thickness uniformity across the whole substrate, measured by high resolution cross-section SEM and HRTEM; **e** cross-section HRTEM image showing the Si/W/UNCD film interfaces, featuring the WC nucleation sites from which the UNCD film grow, while applying the negative voltage on the substrate with respect to the plasma [324].

Diamond Flim Structure	H ₂ flow (sccm)	CH ₄ flow (sccm)	Ar flow (sccm)	Grain size
MCD	200	2	0	3–5 µm
MCD	50	2	50	$\sim 1 \ \mu m$
NCD	30	2	70	100 s nm
NCD	20	2	80	20–50 nm
NCD	15	2	85	10–20 nm
UNCD	10	2	90	3–5 nm

Early research done to investigate HFCVD BEN/ BEG of diamond films were focused on growing mainly MCD to NCD films on high electronic conductivity Si⁺⁺ substrates, mostly on small areas [321, 322], except for work by one group [323], where the authors demonstrated the pathway for HFCVD BEN-BEG of NCD films on up to 100 mm diameter Si substrate, although the film did not cover the Si substrate all the way to the edge (Fig. 4 of Ref. [323]).

Following research on growing NCD films via HFCVD BEN + growth with no bias, systematic research by Auciello's group demonstrated [324] growth of UNCD films via HFCVD BEN-BEG on 100 mm diameter Si wafers coated with nanoscale (60–70 nm thick) W layer, to induce veery dense films, as previously demonstrated for growth of UNCD films on W layers via the MPCVD process [4]. Figure 35c shows the plasma created on the surface of a W-coated Si 100 mm diameter wafer, via application of – 220 V on the substrate, keeping a constant electrical current of 1.5 Amp. through the plasma, producing an extremely uniform UNCD film, in thickness, on a 100 mm diameter Si wafer (Fig. 35d). Cross section HRTEM image (Fig. 35e) shows the formation of WC nucleation sites, induced by the impact of energetic C⁺, CHx⁺ (x = 1, 2, 3) and Ar⁺ ions on the W layer, such that the WC nucleation



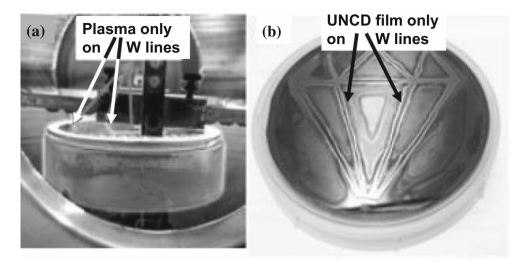


Figure 36 a HFCVD in operation producing UNCD films via BEN-BEG localized on electrically conductive W lines, grown on a Si 100 mm diameter wafer, biased with -200 V with respect to

leads to UNCD film growth (see detailed explanation in [324].

Another key technological development related to the HFCVD BEN-BEG process was the demonstration [324] that this process can be used to grow UNCD films selectively on micro-structured W lines patterned on a 100 mm diameter Si wafer (see Fig. 36).

The selective UNCD film growth process, on patterned W lines, open the pathway to produce selectively grown UNCD layers to subsequently fabricate, in a speedy way, MEMS/NEMS cantilevers to produce MEMS/NEMS-based biosensors and energy generation devices, as described in Sect. 10.3

The UNCD films produced by the MPCVD and HFCVD processes exhibit a unique combination of chemical, mechanical and tribological properties to enable a transformational new generation of MEMS/ NEMS devices, as described in the Sections below.

Integrated CMOS (Driver)/RF-MEMS switches with fast charging/discharging ultrananocrystalline diamond film dielectric

As explained in prior Sections, (RF-MEMS) is one of the important components in Radar and mobile communication electronics. RF-MEMS switches have been demonstrated for GaN-based solid state junction devices [325] and Si-based semiconductors such as Si or SiC [326]. On the other hand, capacitive RF-MEMS switches have been extensively investigated due to their low insertion losses, ultralow power the plasma, to concentrate the C^+ -based species. produced on the heated W filaments, on the W lines; **b** UNCD films grown, selectively, on several biased patterned W lines on the Si wafer.

consumption and ultrahigh linearity. The RF-MEMS capacitive switch involves a flexible metal membrane dangling over a lower metal electrode coated with a dielectric layer. Application of an electric field, between the metal membrane and the lower electrode, results in a physical motion of the membrane towards the electrode, resulting in the change of membrane/electrode capacitance, coupling the RF signal to the desired path. Upon turning the applied electric field off, the membrane is released of the contact to the bottom dielectric layer/electrode (Fig. 37a), interrupting the RF signal. Despite the excellent RF performance of capacitive switches, they have not been used because lack of long-term reliability. Failure modes of RF-MEMS capacitive switches include dielectric breakdown after billions of operation cycles, due to high build-up or electrical charges on electrodes and membranes, resulting in switch failure. An alternative switch failure is induced by stiction of the flexible membrane to the hydrophilic dielectric layer surface (see fig 37a), under increased humidity condition. The most common materials used in the dielectric layer in RF capacitive switches are silicon dioxide (SiO₂) and silicon nitride (Si₃N₄). Under continuous operation, trapped charges in electrodes and membranes induced switch failure. Studies have shown that oxide and nitride dielectric layers charge in 10s of seconds, but they discharge in 100s of seconds, which leads to failure due to the membrane electrostaticinduced stiction, onto the bottom electrode [326]. The

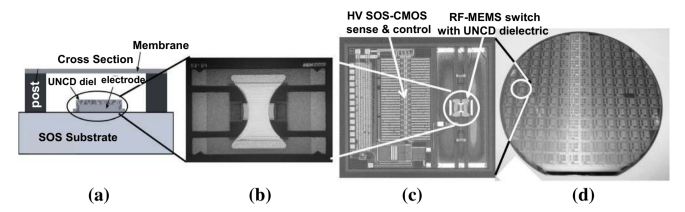


Figure 37 a Cross-section schematic of RF-MEMS capacitive switch with UNCD dielectric layer on metal (W) electrode and metal membrane; b top view optical microscope image of an RF-MEMS switch with metallic membrane and UNCD dielectric layer; c optical picture of a single integrated RF-UNCD/MEMS switch/SOS-CMOS device; d Optical microscope picture of integrated RF-UNCD/MEMS switches/SOS-CMOS devices

RF-MEMS membrane stiction is further increased by the hydrophilic nature of the surface of the conventional metal membranes and oxides and nitrides dielectric layers used today.

The problems described above for conventional RF-MEMS switches were solved by integration of UNCD films, as dielectric layer, into RF-MEMS switches with metal membranes and electrodes (see Fig. 37b–d for details).

Detailed measurements of the dielectric properties of UNCD films in capacitor-type structures and in RF-MEMS switches revealed several insights into physical and device related phenomena, as described below:

- *I*–V measurements indicate that grain boundaries control to large extent the new paradigm charging/discharging behavior exhibited by UNCD films.
- Measurements of the voltage needed to release the switch membrane indicate that this value decreases after charging, which suggests that charging appears to be in the bulk instead of the surface of UNCD [326].
- Measurements of switch release voltage for RF-UNCD-MEMS switches showed that this Voltage decreases monotonically from 13 V to 0 when the charging time is increased from 10 to 500 µs. This rate of voltage' decrease correlates with a time constant of 95 µsec, which are far superior, under comparable fields of 10⁶ V/cm, to the time

fabricated in mass scale on a commercial 200 mm diameter sapphire wafer. Measurements of the response of RF-MEMS switches with UNCD dielectric layer, to electrical excitation, revealed that these switches exhibit time constants that are 5–6 orders of magnitude quicker (about 100 μ s) than those of the switches with oxide or nitride layers (~ 10-100s seconds).

constant (10 s) of RF-MEMS switches with SiO_2 or SiN_x dielectric layers [326].

 Other measurements showed 5–6 orders of magnitude quicker recovery times for RF-UNCD/ MEMS switches than for RF-MEMS switches with SiO₂ or SiN_x dielectric layers. Switches kept in the "on" position for 100 s recovered back to their original condition. In less than 50 µsec. This implies that if switches are cycled off once out of every 100 s, they will be fully recovered from any effects of charging and are ready to be reused anew; thus, the RF-UNCD/MEMS switches have to be turned off only for 0.00005% of the timeline to operate without charging failure.

The R&D performed on developing RF-MEMS with UNCD dielectric layers showed, for the first time, the possibility of operating capacitive MEMS switches almost continuously "on" without an adverse impact on switch reliability.

In conclusion, the data and discussion presented above provide a view of what can be considered as a new paradigm in materials, materials integration, device architecture, and fabrication for a new generation of high performance RF-MEMS switches, featuring UNCD as the new dielectric layer, which may enable a new generation of high frequency systems, using RF-UNCD/MEMS switches as the cornerstone. In addition, Fig. 35d shows that the scaling to growth UNCD dielectric layers, with excellent uniformity on thickness and nanostructure, and dielectric



properties, on 200 mm wafers, and mass scale fabrication of integrated RF-UNCD/MEMS switches/ SOS-CMOS devices [326], provides the bases for commercialization of these new technology.

Integrated multifunctional piezoelectric oxides/ultrananocrystalline diamond (UNCD[™]) films for a new generation of biomedical MEMS sensors and energy generation devices

Powering MEMS-based in-vivo devices is mostly done, currently, via conventional or thin film battery systems. The problem is that current batteries require replacement or recharging more frequently than desired. Therefore, alternative power sources are desirable to replace batteries, which involve undesirably invasive replacement/recharge procedure. For example, defibrillator/pacemakers are a common implantable medical device, which requires an independent power source, operating completely autonomously from the outside world. Currently, defibrillator/pacemaker operation is based on power provided by batteries supplying ~ 0.65 –2.8 A hours for 5 to 9 years [327]. Eventually, the battery for defibrillator/pacemakers needs to be replaced, requiring additional surgery with extra cost and patient discomfort. Power for defibrillators/pacemakers may be provided by an electromagneticbased MEMS generator. Roberts et al. [328] demonstrated an electromagnetic MEMS-based generator, capturing vibrational energy produced by heart muscles, generating power to supplement the pacemaker's internal battery. Initial clinical trials demonstrated that the MEMS energy generator produced up to 17% of the energy required to operate a conventional pacemaker [327]. Further development of the MEMS-based energy generator technology may enable elimination of the costly/invasive surgeries required to maintain the pacemaker, both decreasing medical cost and improving the quality of life for the patient. MEMS-based power generation may also enable implantable biosensors. Two alternative MEMS-based power generation are being explored, namely:

Electrostatic vibration-to-electricity conversion which involves using a MEMS comb drive-type structure, generate electricity from vibrating combs fingers, via change in a vibration-driven capacitance, which causes charge transfer and current flow.

Piezoelectric energy conversion which harvest power from mechanical vibrations of piezoelectric materials. When electric fields are applied to a piezoelectric crystal structure or thin film between two electrode layers, positive and negative ions, in the piezoelectric material, are physically moved in opposite directions, distorting the micro-structure of the crystal, and inducing a polarization in the material, while electrical equilibrium within the crystal is maintained by mobile electrons, creating a current. The effect described above is referred to as the direct piezoelectric effect. On the other hand, the exact opposite phenomenon, the converse piezoelectric effect, is produced by vibration generation in a MEMS oscillating membrane or cantilever-type structure. Piezoelectric generation is frequency dependent, maximized at resonance frequency [329]. The piezoelectric-based MEMS power generator involves using a piezoelectric film encapsulated between two electrode layers to convert the displacement and strain into electricity through the piezoelectric effect (motion of charges induced in the piezoelectric layer). Three materials have been explored as thin films for use in piezoelectric MEMS, namely: lead-zirconatetitanate (PbZr_xTi_{1-x}O₃ (PZT), which main drawback for application to implantable devices is that it contains Pb, making it unacceptable; zinc oxide (ZnO), and aluminum nitride (AlN), the last two being to some extent biocompatible, although substantial research is still necessary to probe that they are biocompatible. More recently, another promising piezoelectric film technology has been demonstrated [330], based on BiFeO₃ (BFO)/SrTiO₃/BiFeO₃ nanolaminate structured films with super-high piezoelectricity compared with PZT, ZrO, and AlN. Initial tests, underway, are demonstrating that BFO may be biocompatible, which is supported by the fact that is formed by Bi (an element in a pharmaceutical drug named Pepto-Bismol), Fe, and O, both in the human body.

A key aspect of integrating a piezoelectric film on an underlying substrate material, to fabricate optimum piezoelectrically actuated MEMS/NEMS biosensor and power generation devices, is to determine which is the best substrate material. In this respect, calculations showed that the resonance frequency/dynamic displacement is largest for the PZT/UNCD hybrid compared to other PZT-based hybrids [331], including Si, nitrides, metals and insulators (see Fig. 38). In addition, UNCD films

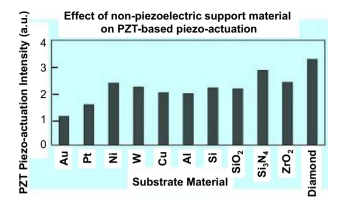


Figure 38 Piezoelectric performance intensity of PZT films grown on different substrate materials for fabrication of piezoelectrically actuated MEMS/NEMS devices, showing the diamond is the best substrate material among all others, currently used in most MEMS/NEMS devices, shown in this figure.

 Table 11 Comparison of mechanical and tribological properties

 between Si, SiC, diamond

Property	Si	SiC	Diamond
Lattice constant (Å)	5.43	4.35	3.57
Cohesive energy (eV)	4.64	6.34	7.36
Young's modulus (GPa)	130	450	1200
Shear modulus (GPa)	80	149	577
Hardness (kg/mm ²)	1000	3500	10,000
Fracture strength (GP)	1.0	5.2	5.3
Flexural strength (MPa)	127.6	670	2944
Friction coefficient	0.4–0.6	0.2–0.5	0.01-0.04

provide the best support material for MEMS resonators and other MEMS/NEMS devices, due to the far superior mechanical and tribological properties shown in Table 11.

Growth and characterization of PZT films on UNCD films for fabrication of piezoelectrically actuated MEMS/NEMS devices

Sol–gel growth process for PZT films involves spreading a solution of Pb acetate tetrahydrate (aldric 99%), Zrpropoxide (aldric 97%) and Ti-isopropoxide, as precursors materials, and 2 methoxyethanol [2MOE] (sigma 99.9%) as a solvent, on the electrode layer grown on the UNCD MEMS/NEMS substrate material, following annealing in air at 450 °C for 5 min to induce the piezoelectric PZT structure (see details in [332]).

RF planar magnetron sputtering-deposition for growth of PZT films can produce PZT films via a relatively simple process, where a plasma us created in front of a stochiometric PZT solid target, inducing bombardment by inert gas ions (mainly Ar ions, the less expensive gas)), which eject atoms (sputtering process) from the surface of the target, with the atoms flowing towards the substrate where they grow the films.Several variations of the RF-sputtering have been developed and are still being optimized for the production of PZT thin films [333]. However, the main problem is that RF magnetron sputtering does no reproduce the stoichiometry of the target on the film, due to preferential sputtering of O atoms, which change the target stoichiometry [333], thus PbO rich targets needs to be used.

Pulsed laser ablation-deposition (PLD) for growth of PZT film this technique involves pulsed laser ablation of solid state stoichiometric PZT target, and a substrate holder opposite to a rotating target at a distance in the range 10–15 cm. In general, a KrF excimer laser (wavelength: 248 nm) is directed through a quartz SUPRASIL II window, into the vacuum chamber, evacuated to ~ 10^{-7} Torr, to evaporate atoms from a stochiometric solid PZT target. The advantage of the PLD technique, over the RF sputtering-deposition process is that the PLD process does not produce change in the stoichiometry of the target, since the laser beams induce evaporation of the material in a stoichiometric way [334, 335].

Metalorganic chemical vapor deposition for growth of *PZT films* This technique uses a bis-tetramethylheptanedionato-Pb tetrakis1-methoxy-2-methyl-2-propoxypoxy-Ti and tetrakis1-methoxy-2-methyl-2-propoxy-Zr, metalorganic mixture of chemical vapor flowing the Pb, Zr. Ti and O atoms on the substrate surface, heated to 450–525 °C. For example, a 70 nm thick *PZT* film grown at 475 °C on an Ir electrode exhibited good ferroelectric (piezoelectric) performance, such as high remnant polarization at 4 V), and small coercive voltage (~0.6 V at 4.5 V) to induce the ions motion inside the material to produce the piezoelectric actuation.

The first attempt at integrating Pt/PZT/Pt/UNCD films into MEMS cantilevers, failed because the UNCD films appeared chemically etched when looking at the bottom Pt/UNCD cross section in a SEM imaging [see review in [332], after growing a PZT film, via the PLD process. Fundamental materials science revealed that O atoms, involved in the



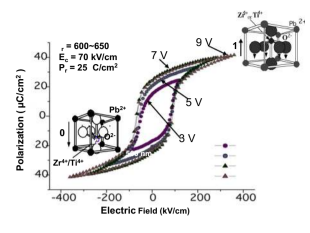


Figure 39 Polarization vs electric field for polarization switching on a PZT film encapsulated between two Pt electrode layers integrated into a UNCD-based cantilever-type MEMS structure for biosensor and power generation.

growth of the PZT films at 450-525 °C, diffused through grain boundaries of the Pt electrode layer in contact with the UNCD film, etching the latter via chemical reaction and formation of volatile CO and CO₂ molecules. This problem was solved by growing, via room temperature sputter-deposition, a layer of TiAl (\sim 30–40 nm) on the UNCD film surface, providing a barrier for O atoms diffusion, based on thermodynamic process, which indicate that oxygen atoms react preferentially with Ti and Al to form stable oxides due to the lowest energy of oxide formation for these elements with respect to all other elements in the periodic table (see details in the review in ref. [332]. The TA barrier inhibits oxygeninduced etching of UNCD (carbon) layer during growth of PZT at relatively high temperature (450–500 °C) in oxygen. PZT films were grown by the PLD technique described above.

Measurement of the polarization vs. applied voltage, between top and bottom Pt electrodes encapsulating the PZT film, inserted in cantilevertype MEMS structure for biosensors and power generation devices, showed excellent polarization saturation (see Fig. 39).

Fabrication of integrated Pt/PZT/Pt/UNCD heterostructure MEMS cantilevers

For the R&D described in this Section, the fabrication of Integrated Pt/PZT/Pt/TiAl/ UNCD heterostructure films MEMS/NEMS *c*antilever-based biosensors and power generation devices involved the following steps (see details in [332]):

- Growth of 1 μm thick UNCD layer on Si (100) substrate coated with 1 μm SiO₂ sacrificial layer
- (2) Growth of a 20 nm thick TiAl barrier layer on the UNCD film to stop O atoms diffusion, during growth of PZT films in O gas at ~ 450–525 °C, which would etch the C-based UNCD film;
- (3) Growth of 180 nm thick Pt layer (bottom electrode) on top of the TiAl barrier layer;
- (4) Growth of 60 nm thick PbZr_{0.47}Ti_{0.53}O₃ piezoelectric layer, via PLD at 550 °C, in 100 mTorr of O gas;
- (5) Growth of the top 50 nm thick Pt layer (top electrode) to complete the capacitor-like structure to measure the polarization properties (see Fig. 39) of the PZT layers integrated with the UNCD films, and piezo-actuation via voltage application between the top and bottom Pt electrode layers in the Pt/PZT/Pt capacitor-type structure produced on the same UNCD film used for MEMS cantilevers; Steps 1 to 5 are shown in Fig. 40a.
- (6) Fabrication of UNCD (~ 140 μm long × 1–10 μm wide) cantilevers, using a Focused Ion,

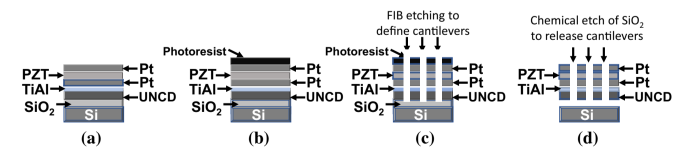
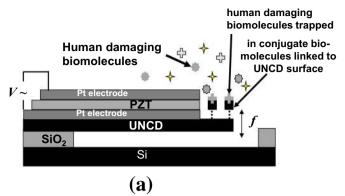


Figure 40 a Integrated Pt/PZT/Pt/TiAl/IUNCD structured films grown on SiO₂/Si substrate; **b** Photoresist grown on top Pt electrode layer; **c** Focused Ion Beam etching, through photoresist,

to define cantilevers beams; **d** chemical all etching of the SiO_2 layer to release the integrated Pt/PZT/Pt/TiAl/IUNCD cantilevers.



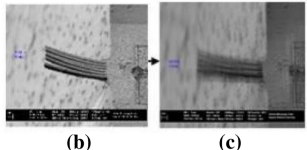


Figure 41 a Schematic of integrated Pt/PZT/Pt/UNCD heterostructure MEMS cantilevers-based biosensors, developed to sense anthrax molecules; **b** MEMS image of an array of Pt/PZT/Pt/UNCD heterostructure MEMS cantilevers fabricated as explained in the text above this figure; **c** SEM video image of

Beam (FIB) system with a 30 keV Ga⁺ ion beam to etch the integrated Pt/PZT/Pt/TiAl/UNCD films to define the cantilevers (see Figs. 40b,c),

(7) Chemical etching of the SiO₂ layer, using HF acid, which etches only the SiO₂ layer, without any chemical attack on UNCD, thus liberating the cantilever formed by the integrated Pt/PZT/Pt/TiAl/UNCD films (Fig. 40d).

Integrated Pt/PZT/Pt/TiAl/IUNCD film-based MEMS cantilevers can be processed also via lithography and RIE processes, as demonstrated in prior R&D [332].

Demonstration of piezoelectrically actuated Pt/PZT/Pt/ UNCD film-based heterostructure MEMS cantileversbased biosensors

Figure 41 shows the demonstration of a biosensor to detect deadly anthrax molecules, via chemical growth, on the surface of the exposed UNCD end of the cantilever, biomolecules that can react only with anthrax molecules (see Fig. 41a). Figure 41b shows an SEM image of the integrated Pt/PZT/Pt/UNCD heterostructure MEMS cantilevers-based biosensors actuated via application of 3 V AC (1 Hz to 1 MHz) between the top and bottom Pt electrodes, inducing polarization of the piezoelectric layer between them, which made the cantilevers vibrate for up to 2 billion cycles without any degradation (Fig. 41c). The biosensing of anthrax molecules happen when those molecules are captured by the conjugated

Pt/PZT/Pt/UNCD heterostructure MEMS cantilevers excited by 3 V applied between the top and bottom Pt electrodes, encapsulating the piezoelectric PZT film, for up to 2 billion cycles without degradation.

biomolecules grown on the UNCD cantilever's surface (see Fig. 41a) and the cantilever vibration is changed.

The piezoelectrically actuated Pt/PZT/Pt/UNCD heterostructure MEMS cantilevers-based biosensors demonstrated for detection of anthrax can be applied to detection of other biomolecules by growing the appropriate conjugate biomolecules on the surface of the UNCD layer.

Demonstration of piezoelectrically actuated Pt/PZT/Pt/ UNCD heterostructure MEMS cantilevers-based power generation device

Cardiac cell mediated actuation of piezoelectrically actuated UNCD-based MEMS biopower generation device is based on myocardium' cells exhibiting compression and relaxation cycle beating in the 2–3 Hz range, autonomously, when properly cultured. Cardiac cells from certain animals, such as birds, actuate even at higher frequencies. The actuation is driven by the energy supplied to the cells by nutrients in the fluid environment where the cells live.

Based on the above information, the opportunity exists for investigating the integration of myocardium cells with piezoelectrically actuated cantilevers to achieve biopower generation via cell beating-induced mechanical vibration of the cantilever and generation of electrical potential from the mechanical deformation of the piezoelectric layer,

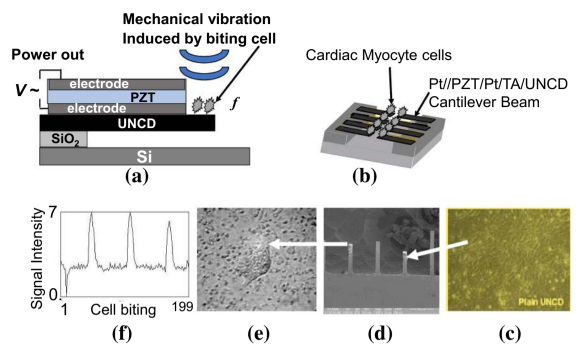


Figure 42 a schematic of a MEMS power generation device based on integrated Pt/PZT/Pt/ TiAl/UNCD heterostructure filmbased MEMS cantilever; b schematic of array of integrated Pt/ PZT/Pt/ TiAl/UNCD cantilevers, with heart cells grown on the exposed surface of the UNCD layer to induce the mechanical vibration, generating power (~ 3 mW already demonstrated by Auciello's group [337]); c SEM image of cells of mice heart grown

which supplies electrical through top and bottom film electrodes encapsulating the piezoelectric layer.

Cardiac cells can be grown on the end of the exposed UNCD cantilever surface of the integrated Pt/PZT/Pt/UNCD cantilever, and hopefully in the near future replacing the non-biocompatible Pbbased PZT by BiFeO₃, a new alternative biocompatible (made of biocompatible Bi, Fe, and O) piezoelectric material under investigation [330], exhibiting higher polarization than PZT. The growth of different types of biological cells on the surface of UNCD films has been recently demonstrated [336, 337]. The beating cardiac cell makes the cantilever vibrate, and the strain produced on the piezoelectric layer, induced by Pb²⁺, Zr⁴⁺, and Ti⁴⁺ ions displaced in one direction and the O^{2-} ion in the opposite direction ((inverse piezoelectric effect), thus inducing the PZT layer sandwiched between bottom and top electrode layers to induce voltage pulses extracted between the electrodes. The part of the cantilever containing the cells can be immersed in blood cell fluid in the human heart, which provides the fluid with nutrients that energize the cardiac cells, enabling the on the exposed UNCD surface (see **a**) of the cantilever; **d** low magnification SEM imaging showing array of Pt/PZT/Pt/ TiAl/ UNCD cantilevers, with mice hearts' cells grown on the end; **e** higher magnification SEM image of a mice's heart cell grown on the surface of a UNCD cantilever; (f) signal showing the biting pattern of the mice heart's cell shown in **e**.

piezoelectric actuated power generation MEMS device as shown in Fig. 42.

The integrated Pt/PZT/Pt/TiAl/UNCD cantilever-type MEMS power generation device powered by heart cells, may enable replacement of the Liion batteries used today to power defibrillator/pacemakers, which need to be replaced in undesirable short periods of time (6–9 years from the time of first implantation [338]), which requires expensive replacement cost and patient discomfort [338].

The information presented in "Integrated multifunctional piezoelectric oxides/ultrananocrystalline diamond (UNCD[™]) films for a new generation of biomedical MEMS sensors and energy generation devices", secion including all sub-Sections, above shows that the integrated biocompatible Pt/piezoelectric/Pt/TiAl/UNCD MEMS/NEMS cantileverbased devices may provide transformational technologies for new generation of biosensors and biologically based power generation devices to make a revolutionary impact in improving the quality of life of people worldwide.

Demonstration of piezoelectrically actuated Pt/AlN/Pt/UNCD heterostructure membranes for MEMS drug delivery devices

Growth of AlN films on integrated Pt/Ti adhesion layer/UNCD film

AlN films for piezoelectrically actuated MEMS devices are synthesized mainly in the Wurtzite hexagonal crystallographic structure with c-axis (002) orientation perpendicular to the surface (XRD diffraction peak appears at 36°), corresponding to the crystal orientation that yields the highest piezoelectric constant in addition to a high acoustic wave velocity [339, 340]. AlN (002) oriented films were grown, using reactive sputter deposition (AJA ATC Orion system), involving sputtering of Al atoms from a metallic target, via Ar ions bombardment from an Arplasma, and flowing N₂ gas to produce the AlN films on Pt electrode layers, at 500 °C and pressure of 3mTorr. AlN layers with thickness in the 260-420 nm range were grown on Pt/Ti bottom electrode layer grown on UNCD films on a 100 mm diameter Si wafer. XRD analysis revealed a diffraction peak at 36.05°, corresponding to the hexagonal AlN (002) orientation. which induces the highest piezoelectric coefficient in AlN films (see detailed information in [340]), Research showed that AlN films grown on chemical mechanically polished (CMP) B-doped UNCD (B-UNCD) films with Pt electrode on the surface exhibit surface roughness of about 0.2 nm rms and high (002) orientation, confirmed by XRD analysis [340], similar to UNCD films grown on atomically flat semiconductor surfaces. In addition, measurement of the piezoelectric coefficient revealed that the extremely smooth (002) oriented AlN films exhibit 5.3 pm/V piezoelectric coefficient [340], which is the highest among currently reported values for AlN films. The results presented here indicate that the smoothness of substrate surface is critical to achieve highly (002) oriented very thin AlN films.

Fabrication of integrated Pt/AlN/Pt/Ti/ UNCD film-based piezoelectrically actuated MEMS drug delivery device

The fabrication of Pt/AlN/Pt/Ti/UNCD film-based piezoelectrically actuated drug delivery membranes involved the following steps: (1) UNCD thin films were grown by the MPCVD process, using the parameters described in Sect. 10.1, on Si substrates after a nano-diamond seeding process on the silicon surface, (2) AlN/Pt/Ti (adhesion layer on UNCD) films were grown on to the UNCD layer (see Fig. 43a) [340]; (3) photolithography, using a specially designed mask, was used to define a window on the backside of the Si wafer, (4) reactive ion etching was used to pattern the Si₃N₄ mask to subsequently produce chemical etching of the Si substrate to produce the cavity sustaining the UNCD drug deliver membrane on the front side of the wafer (see Fig. 43b). Square membranes of UNCD, with lateral dimensions in the range 200–100 µm, with thickness in the range 100-500 nm, were fabricated and characterized by Raman spectroscopy, optical microscopy, scanning electron microscopy (SEM) (see Fig. 43) and reflectometry.

Pt/AIN/Pt/Ti UNCD piezo-actuated drug delivery membrane

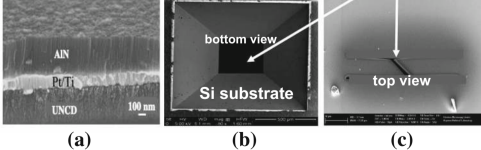


Figure 43 a SEM image of AlN/Pt/Ti (adhesion layer)/UNCD films grown on a Si wafer; **b** patterned etched Si cavity showing the bottom surface (UNCD) of the integrated Pt/AlN/Pt/Ti/UNCD

piezoelectrically actuated drug delivery membrane; c top view SEM of the integrated Pt/AlN/Pt/Ti/UNCD piezoelectrically actuated drug delivery membrane.



The active drug delivery device involves the actuation of a piezoelectric valve involving an AlN film, as the biocompatible piezoelectric film encapsulated between two Pt electrodes and the whole structure grown on an UNCD film grown on a Si wafer. This device is capable of controlling drug delivery dose with an applied electric field between the two Pt electrodes encapsulating the piezoelectric AlN layer, which actuate the lever valve, shown in Fig. 43 above, inducing the flow of drug containing fluid from a reservoir to the outside world.

The morphology and operation of the piezoelectrically actuated cantilever is shown in the image from a numerical simulation (Fig. 44), where the piezoelectric layer is grown onto the top of the UNCD membrane, occupying a small part of the latter thus increasing the strain of the membrane on the valve.

The R&D described in this Section demonstrated that the integration of two unique dissimilar materials, such as a piezoelectric ALN and a diamond based

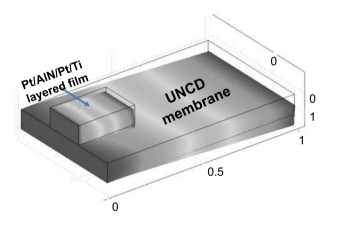


Figure 44 Computer simulation showing the morphology and operation of a piezoelectrically actuated Pt/AlN/Pt/Ti/UNCD cantilever valve for an active drug delivery device.

UNCD film can produce transformational new generation of external and implantable MEMS/NEMS sensors and drug delivery devices to improve the quality of life of people worldwide.

Fabrication and characterization of MEMS structures for electron field emission devices based on N-UNCD-coated Si surface

Carbon nanotubes and NCD have been investigated for the last two decades for fabrication of electron field emission devices, because their low power consumption and potential for miniaturization. Long life electron field emission materials are extremely powerful for long life electron emission sources for mass spectrometers for space exploration where electron sources are exposed to harsh environments, such as miniaturized mass spectrometers for use as in situ chemical analyzers on the moon and other planetary surfaces, long-lived electron source, to generate ions from gaseous sample using electron impact ionization. In relation to the use of UNCD film technology, R&D performed several years ago demonstrated that the MPCVD process discussed in Sect. 10.1, involving flowing CH_4/N_2 gas mixture into a vacuum chamber, coupling microwave power, creating a plasma involving C⁺, CHx⁺ (x = 1, 2, 3) and N⁺ ions and associated neutrals, produced the growth of UNCD films with N atoms incorporated in the grain boundaries, defining these films as N-UNCD [4, 341], which have been demonstrated to exhibit superb electron field emission properties.

N-UNCD films were grown on a range of silicon substrates with varying microstructure, such as sharpened tips (Fig. 45a), elongated ridges

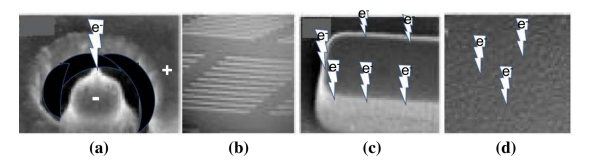
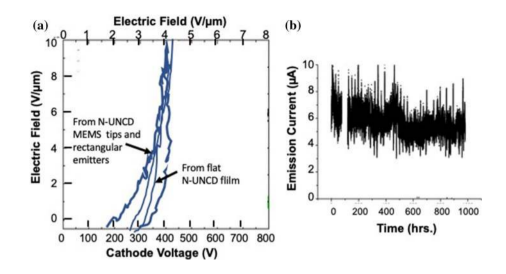


Figure 45 a Single N-UNCD-coated MEMS tip/open electrode structure in a large array for field emission device; \mathbf{b} low and \mathbf{c} high magnification SEM images of ridge electron emission structures; \mathbf{d} flat N-UNCD electron emission structure.

Figure 46 a Electric Field vs Cathode Voltage (Electric Field) for electron emission from N-UNCD-coated Si tips and ridges and from flat N-UNCD surface; b electron emission current vs time for a N-UNCD-coated Si tip array.



(Figs. 45b,c), and planar surfaces (Fig. 45d). The current–voltage characteristics were measured for each topography in high vacuum ($\sim 10^{-7}$ Torr). Lifetime tests were also performed for a representative geometry in high vacuum ($\sim 10^{-8}$ Torr) to determine the viability of N-UNCD cold cathodes for use in long-duration space missions and other devices used in the Earth environment.

N-UNCD-coated Si cathodes based on MEMS tips and ridges structures (Fig. 45a-c) and flat N-UNCD cathode (Fig. 45d), all exhibited electric field emission of ~ $3 \text{ V/}\mu\text{m}$ (Fig. 46a), which is one of the lowest among field emission devices demonstrated today [342]. The explanation for emission field from flat N-UNCD surface similar to emission from tips and ridges is that the electron emission is coming from grain boundaries of the N-UNCD films, which are $\sim 1-2$ nm wide, concentrating voltages very efficiently to emit electrons at low electric fields, and the grain boundaries emission also dominate emission from tips and edges of ridges, which have much larger dimensions (≥ 100 nm) that the grain boundaries. Another key result from the electron field emission measurement from a N-UNCD-coated tip array is that they exhibited tens of microamps of emission current with little degradation for 1000 h (Fig. 46b) (the longest stable electron field emission demonstrated today from a field emission device).

Summarizing, electron field emission from N-UNCD-coated Si tip and ridge arrays and flat N-UNCD surfaces shows that N-UNCD films provide the basis for superior electron emission devices for multiple application in outer space and on Earth.

Conclusions

Based on the information presented in this review, the outlook for Microcrystalline Diamond (MCD), Nanocrystalline Diamond (NCD) and (UNCD) Ultrananocrystalline Diamond -based MEMS/NEMS technologies is promising from various points of view, namely: the superior mechanical, tribological, chemical corrosion resistant, electronic, and biocompatible properties demonstrated for diamond-based MEMS/NEMS devices provides the pathway for the development and marketing of new technologies using the superior diamond-based MEMS/NEMS devices, over for example Si-based MEMS impact sensors used in deployment of safety bags in cars, when in a high impact accident, MEMS devices measuring tires pressure, MEMS devices used in airplanes, external and implantable medical MEMS/NEMS sensors and drug delivery devices, and in many other applications.

Acknowledgements

O. Auciello acknowledges the support from the University of Texas at Dallas through his Distinguished Endowed Chair Professor position (2012present) and the funding from DOE and DARPA, when working at Argonne National Laboratory, doing R&D for developing UNCD-based MEMS/ NEMS (1996-2012). D.M. Aslam acknowledges the support provided by NSF Engineering Research Center (ERC) for Wireless Integrated Micro Systems (WIMS) during 2000–2010 awarded to University of Michigan, Michigan State University and Michigan Technological University. O. Auciello and D.M. Aslam are thankful to Dr. Zongliang Cao for providing some early written materials, over 10 years ago, that ultimately led to this detailed review. O. Auciello and D.M. Aslam acknowledge the great contributions of the many scientists, postdocs, and students, who contributed to the science and technology development of MEMS/NEMS devices/systems through many years, as indicated by the many references provided in this review article.

Compliance with ethical standards

Conflict of interest The authors declare that they have no conflict of interest.

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