Nitrogen modification of hydrogenated amorphous carbon films

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The effect of nitrogen addition on the structural and electronic properties of hydrogenated amorphous carbon films has been characterized in terms of its composition, sp³ bonding fraction, infrared and Raman spectra, optical band gap, conductivity, and paramagnetic defect. The variation of conductivity with nitrogen content suggests that N acts as a weak donor, with the conductivity first decreasing and then increasing as the Fermi level moves up in the band gap. Compensated behavior is found at about 7 at. % N, for the deposition conditions used here, where a number of properties show extreme behavior. The paramagnetic defect density and the Urbach tailwidth are each found to decrease with increasing N content. It is unusual to find alloy additions decreasing disorder in this manner. © 1997 American Institute of Physics.

I. INTRODUCTION

Recent studies of nitrogen additions to amorphous carbon (a-C) have been motivated by two factors; the synthesis of the possible superhard compound C₃N₄,¹ and the electronic doping of a-C. We are interested here in the semiconducting forms of a-C and hydrogenated a-C (a-C:H) which can be doped electronically using nitrogen. Meyerson and Smith² observed that a-C:H could be doped n- and p-type by phosphorus and boron, respectively, in a similar fashion to a-Si:H. They concluded that there was true substitutional doping by observing the thermopower sign reversal expected as the conductivity changed from n-type to p-type.³ However, the doping effect was weaker than in a-Si:H. Indeed, Jones and Stewart⁴ found still lower doping efficiencies and they questioned whether true doping was occurring. Interestingly, they observed that nitrogen was able to increase the conductivity of a-C:H, and this has since been repeated by numerous other groups.⁴−¹⁰ However, it has been unclear whether N increased the conductivity by doping—raising the Fermi level towards the conduction band, or by graphitization of the bonding—narrowing the band gap.⁷

Recently, a highly sp³ bonded form of a-C denoted tetrahedral amorphous carbon (ta-C) has been deposited by filtered cathodic vacuum arc (FCVA).¹¹ Ta-C is a semiconductor with a 2.5 eV wide band gap. Undoped ta-C is moderately p-type.¹² Ta-C can be doped in a rather uncontrollable fashion by P.¹³ It can be doped more controllably by N,¹⁴,¹⁵ and N was found to first decrease the conductivity, then increase it, as E_F moves upwards in the band gap, from near the valence band, through midgap, to close to the conduction band edge E_C. The optical gap and the fraction of sp³ bonding were both found to remain constant during these N additions, and they only decreased at higher N contents. It was clear that N acted as a substitutional donor in ta-C, although with low doping efficiency.

Subsequently, a hydrogenated analogue of ta-C, ta-C:H, was prepared by deposition from a low pressure plasma beam source.¹⁶ Ta-C:H has an optical gap of 2.7 eV, but again the conductivity activation energy for undoped ta-C:H is much less than half the optical gap, suggesting that E_F lies well away from midgap. N additions to ta-C:H are also found to give a strong increase in the conductivity, while the optical gap remains unchanged.¹⁷ In this case, it was not possible to observe the initial decrease in conductivity as E_F passed through midgap. These latter results, therefore, suggest that
unpaired and available for doping. A variant of this (c), forms a trivalent C site and transfers the unpaired electron to the C, to give a positive N⁺/C⁻ defect pair. This group has all electrons paired, like the analogous group in a-Si:H.²⁶ There are other configurations, because N can also form π bonds. N can substitute for C in a benzene ring [(d)/(e)]. Nitrogen can have just two neighbors within the ring, using three bonding electrons (2 σ and one π), leaving two electrons as a lone pair, Fig. 1(d). This pyrridine configuration is nondoping. Alternatively (e), three can form σ bonds, a fourth forms a π bond (as in benzene), and the fifth is unpaired and enters an antibonding π* state. This electron is available for “π’” doping. Another alternative (f) is for the N to have to bond to three neighbors, as in (e), but in a five-fold ring.²⁴ so it uses three electrons in bonds and two as a lone pair. This “pyrrole” configuration is also nondoping. There are also olefinic variants of π bonding. A double bonded unit (g) uses two electrons in σ bonds, one in a π bond, leaving a lone pair. This is nondoping. A double bonded unit (h) uses three electrons in σ bonds, one in a π bond, and the fifth in a nonbonding π* state, available for doping. The last configuration is a “chain terminating” cyano (or nitrite) grouping (i), in which N forms a triple bond and a lone pair. This in nondoping. It is relatively stable, being related to that found in the N₂ molecule itself. This variety of configurations available for N, most of which are nondoping, indicates why N is likely to be a weak dopant and why the behavior of N in a-C:H:N needs careful analysis.

II. EXPERIMENT

The nitrogenated a-C:H (a-C:H:N) films were deposited in a magnetically confined rf plasma enhanced chemical vapor deposition (PECVD) system.¹⁹ The substrate was placed on the rf powered electrode which acquires a negative dc self-bias voltage. This causes an ion bombardment of the growing film which is believed to be responsible for creating the sp³ bonding. An encased permanent rare earth magnet (Nd-B-Fe) is used around the substrates to increase the plasma density. The deposition was carried out at a fixed pressure of 0.1 Torr from a CH₄, He, and N₂ gas mixture, with flow rates of 5 sccm for CH₄, 45 sccm for He, and 0–20 sccm for N₂. The magnetic field and the He increases the ionization of the methane plasma, giving a plasma electron temperature of over 6 eV, which results in a higher deposition rate and a lower bias voltage for a given rf power. The substrate temperature was maintained at 20 °C, although the surface temperature was seen to vary from 30–100 °C depending on the rf power. The a-C:H:N films were deposited on [100] Si and quartz substrates.

The composition of the a-C:H:N films was determined by a combination of elastic recoil detection analysis (ERDA), Rutherford backscattering (RBS), and nuclear reaction analysis (NRA). The N content was found using the \(^{15}\text{N}(d,p)^{15}\text{N}\) nuclear reaction. Infrared (ir) spectra were collected at room temperature on a Mattson FTIR spectrometer. Raman spectroscopy measurements were conducted in the back scattering mode using an Ar laser (wavelength 488 nm) operating at a power of 300 mW. The scattered radiation was
analyzed with a 1 m focal length double-pass monochromator (Yobin-Yvon, Ramanor model HG2-S) equipped with holographic gratings (2000 lines per mm), and was detected by a cooled (−35 °C) GaAs photomultiplier tube (RCA, Model C31034A-02), operated in photon counting mode. The monochromator stokes Raman spectra were recorded between 500 and 3000 cm⁻¹ and stored in a multichannel analyzer. Optical absorption spectra were derived from transmission and reflection measurements on a Philips uv-VIS spectrometer in the wavelength range 200–900 nm. The conductivity was measured using metal-carbon-metal sandwich structures using Au and Al electrodes evaporated onto the samples at pressures better than 10⁻⁶ Torr. Electron spin resonance (ESR) studies were performed on a Bruker EPR machine such that the signal did not saturate. The microwave power was varied between 200 μW and 2 mW using a microwave frequency of 9.3 GHz. The modulation amplitude was 0.9 G using a modulation frequency of 100 KHz.

Transmission electron microscopy (TEM) samples were prepared by dissolving the (100) Si substrate in a HF:HNO₃ deionized water solution and releasing the a-C:H:N samples. These films were then floated onto Cu grids. Electron energy loss spectra (EELS) were measured on a VG501 scanning transmission electron microscope (STEM) operated at 100 KeV. The spectra were acquired on a dedicated parallel EELS detector with a collection angle of 10 mrad and an energy resolution of 0.6 eV. The specimen thickness was kept constant (60–80 Å) with the EELS spectra collected over an area of approximately 100 nm diam. The collected data was corrected for multiple scattering using the log ratio method. Low loss spectra were also measured to derive the valence plasmon energy and the joint density of states. Typical collection times for the low loss region was then analyzed in terms of dielectric function. This allowed the extraction of the real and imaginary parts of the dielectric function. This allowed the extraction of the optical conductivity or joint density of states. This allowed the extraction of the real and imaginary parts of the dielectric function. This allowed the extraction of the optical conductivity or joint density of states. This allowed the extraction of the real and imaginary parts of the dielectric function. This allowed the extraction of the optical conductivity or joint density of states.

III. RESULTS

A. Composition and growth

The variation of H and N content of the a-C:H films as a function of nitrogen gas flow ratio, \( R = [N_2]/[CH_4] \) is shown in Fig. 2. The N content is seen to increase quite rapidly up to \( R = 1 \), and then increase more gradually. The data shows that the incorporation ratio is much less than one, presumably because N is less easily ionized than CH₄. Although the ionization potential of N₂ is 11.1 eV, in order to get N⁻ ions an energy of 14.5 eV is required. To generate CH₄⁺ and CH₃⁺ species from the CH₄ molecule, ionization energies of 12.6 and 14.3 eV are required. The H content of undoped a-C:H is 21 at %. This value decreases gradually with N incorporation, up to a gas flow ratio of 1.0, and it then falls more rapidly.

The growth rate depends strongly on N addition, and it appears to decrease in a roughly exponential fashion until an atomic N content of 3%, as seen in Fig. 3(a). After this, the decrease in growth rate is more gradual.

Compressive stress is an intrinsic feature of films grown here under conditions of ion bombardment. The compressive stress was derived using the method proposed by Argon et al. based on the change in curvature of the substrate using Stoney’s equation. The variation of the stress as a function of atomic N content is shown in Fig. 3(b). The stress is seen to decrease with N content. The lower stress at higher N contents is desirable, as it allows thicker, adherent films to be deposited. The maximum thickness increases from about 400 nm for a-C:H (non-nitrogenated) to over 1 μm for 15 at. % N content.

B. Transmission electron microscopy and electron energy loss spectroscopy

The TEM studies of the a-C:H:N films show them to be completely amorphous with no crystalline inclusions. Two diffused rings centred at 2.1 and 1.1 Å are seen in diffraction mode which correspond to amorphous carbon. The structure of the film changes dramatically when the N content
exceeds a threshold of 14 at.%. At low N content, the films are extremely smooth, with peak-to-peak roughness of less than 5 nm over atomic force microscopy (AFM) scans of 1 μm. For N contents over 14%, the surface becomes rougher, with amorphous nodulelike surface features with average heights of 20 nm and diameters of 50 nm.

The nature of the C and N bonding was derived from EELS. The spectrum consists of absorption edges for each element's core level, and the absorption just above the edge is proportional to the local density of conduction states about that atom. The carbon K edge consists of a step at 290 eV due to transitions from the C 1s core level to σ* states and a peak at 285 eV due to transitions to π* states of sp² sites (and sp¹ sites). The N K edge spectrum appears around 400 eV and consists of a similar step and peak features, Fig. 4. The C sp² fraction can be deduced from the C K edge, as the ratio of the area of the 285 eV peak to the 290 eV step, normalized to its value in fully sp² graphitic C. This same ratio is then assumed to apply to that for sp² N sites. The EELS spectrum could also be used to derive the N/C composition.

The variation of sp² fraction for C and N sites as a function of N content is shown in Figs. 5(a) and 5(b). The C sp³ fraction is seen to remain within the range 30–40% up to a N content of 7 at.%, where the sp³ fraction reaches a maximum of 45%, and then it decreases gradually to 20% sp³ content at 14 at.% N. The latter fall corresponds to graphitization which usually occurs at high N content. The sp² fraction at N sites varies more dramatically. At low N content, the N sp¹ fraction is low. It then rises sharply to about 80% of N sites at 7% N, and it then decreases again to about 40% at higher N contents. This result is one of a number to indicate that 7% N is a critical composition.

The valence plasmon energy was also measured by EELS, and used to derive a value for the mass density. The plasmon is 22.8±0.3 eV across the range of N contents.

C. Infrared spectroscopy

The local bonding of C, H, and N sites can be deduced from a representative infrared absorption spectrum as shown in Fig. 6. The spectra can be divided into four regions, the C–C, C–H and C–N modes from 0 to 1700 cm⁻¹, C═N stretching modes from 2065 to 2260 cm⁻¹, C–H stretching modes from 2700 to 3100 cm⁻¹, and N–H stretching modes from 3150 to 3400 cm⁻¹, according to assignments in molecules.

The variation of the intensity of these modes as a function of N content is shown in Fig. 7. The intensities of the C–H stretching modes are seen to decline rather rapidly with increasing N content. These modes are assigned to vibrations of sp³ and sp² sites. Notice that sp¹ sites would introduce modes around 3300 cm⁻¹. The finite value of modes around 3300 cm⁻¹ in N-free films indicates a contribution from H–C=C- groups as well as =N–H groups. Infrared peaks observed at 2325 cm⁻¹ in cathodic arc deposited a-C:H:N films by Merchant et al. has also been ascribed to triple bonded carbon.

FIG. 4. Typical EELS spectrum for a-C:H:N with the assignment of prominent bands.

FIG. 5. (a) Carbon and (b) nitrogen sp² fraction of a-C:H:N films vs N content.

FIG. 6. Typical infrared spectrum for a-C:H:N with the assignments of prominent bands in an a-C:H:N film with an atomic N content of 11%.
The intensity of the N–H modes at 3140–3350 cm$^{-1}$ is seen to increase rapidly with N content. Similar modes ascribed to N–H modes in the data of Kaufman et al. can be associated with an increase in N–H bonds.

The modes below 1700 cm$^{-1}$ are particularly interesting. The C–C stretching modes lie up to about 1350 cm$^{-1}$ for sp$^3$ C, up to 1600 cm$^{-1}$ for aromatic sp$^2$ C and up to 1700 cm$^{-1}$ for olefinic sp$^2$ C. The $\text{NH}_2$ (1560–1580 cm$^{-1}$), $\text{NH}_3$ (1640–1675 cm$^{-1}$), $\text{C}=\text{N}$ (1600–1700 cm$^{-1}$), $\text{C}=$ (1675–1700 cm$^{-1}$) groups also make a contribution to these modes, but is unlikely to be of significant magnitude especially at low N contents. In a-C:H the C–C modes are not Raman active (only Raman active), because the C–C bond is not polar. However, in a-C:H:N, the introduction of polar C–N bonds may render these modes Raman active. Moreover, as the modes are delocalized, particularly around aromatic rings, the N atoms tend to make all modes to an extent Raman active. For example, the N atom in the pyrridine ring, Fig. 1(d), will make all the modes of a distorted ring Raman active. Figure 7 shows that the intensity of the 1500–1785 cm$^{-1}$ modes, which is one of the dominant peaks in this region, first increases and then decreases with increasing N content. The increase corresponds to the increasing oscillator strength of mixed C–C–N modes of both aromatic and olefinic units. The decrease presumably corresponds to a fall in the concentration of aromatic and olefinic groups due to an increase in cyano (nitrile) groups at high N contents. The offset observed in the data confirms the fact that the C–C stretching modes dominate initially with any consequent changes arising from the C–C–N modes.

Modes at 2065–2260 cm$^{-1}$ correspond to the terminating cyano (nitrile) group $\text{C}=$ (N), Fig. 1(i), the intensity of which is seen to rise rapidly with increasing N content in Fig. 7. There are no corresponding modes of C atoms in this range. Schwan et al. also found a rapid increase in the C=N and C≡N bonded modes in their plasma deposited a-C:H(N) films with increasing N content in the source gases.

### D. Raman spectroscopy

Raman spectroscopy is a widely used technique to characterize amorphous and noncrystalline carbons. The Raman spectrum of a noncrystalline C in general consists of two features, a $G$ peak at around 1550 cm$^{-1}$ due to optical zone center vibrations ($E_{2g}$ mode) in graphite of aromatic rings, and a $D$ peak or shoulder around 1360 cm$^{-1}$, due to the disorder activated optical zone edge modes of microcrystalline graphitic sheets. The modes of sp$^3$ bonding in a-C tend not to show up in Raman, because of their low scattering cross-section. Tamor and Vassel have provided a useful empirical correlation of the position, width, and intensity ratio of the $G$ and $D$ peaks to the properties of a-C:H. The Raman spectra of our films were quantitatively analyzed using a computer code in order to reproduce the data as a sum of two Gaussians. The position, width, and intensity of each line were assumed as fitting parameters and the least square fit of the experimental data was attempted. Figure 8 shows that the position of the $G$ peak, $\omega_G$, increases with N content. This generally corresponds to increasing graphitic character in a-C:H.3,36 The width of the $G$ peak $\Gamma_G$ decreases with increasing N content. $\Gamma_G$ is a measure of the network disorder, as it is in a-Si, and its reduction is consistent with the fall in stress with N content seen in Fig. 3(b). The position of the $G$ band shifts to higher frequency and its bandwidth decreases. These results show that N incorporation in the a-C:H films favor the formation of sp$^2$ units according to the Raman data. These microstructural modifications (and decrease in the average coordination number) may account for the observed reduction in the internal stress of these films.

### E. Optical properties

The variation of the optical (Tauc) gap as a function of N content is shown in Fig. 9(a). The optical gap is seen to increase from 1.65 eV in a-C:H itself up to 2.10 eV at 7% N. The gap then decreases slightly as the N content rises from 7% to 14%. The latter decline is presumably due to the inter-
increased sp\textsuperscript{2} fraction induced by the addition of N, Fig. 5(a). The local maximum in the gap at 7\% N is further indication of a critical composition.

The refractive index shown in Fig. 9 generally declines with increasing N content. The refractive index is related to the average of ‘Penn’ band gap $E_g$ and the plasmon energy $E_p$ by the Penn model,\textsuperscript{38}

$$n^2 - 1 = \varepsilon_0 - 1 = E_p^2/E_g^2.$$ \hspace{1cm} (1)

It has already been noted that the plasmon energy remains constant at 22.8 eV with N content, according to the low loss EELS. Thus, the Penn gap increases with increasing N content. The average gap corresponds to the average bonding antibonding splitting and is much larger than the minimum optical gap. The average gap of a-C:H from Eq. 1 is 11.3 eV. This is less than for diamond, 13.6 eV, because of the presence of $\pi$ states which have a narrower gap. The average band gap is expected to increase with increasing N content. The average gap corresponds to the average bonding antibonding splitting and is much larger than the minimum optical gap. The average gap of a-C:H from Eq. 1 is 11.3 eV. This is less than for diamond, 13.6 eV, because of the presence of $\pi$ states which have a narrower gap. The average band gap is expected to increase with increasing N content.

The width of the optical absorption edge shown in Fig. 9(c), the Urbach energy $E_U$, declines with increasing N content up to 7\% and then remains relatively constant. The Urbach energy is a measure of the disorder of the atomic network. This decrease suggests that the disorder of a-C:H:N may reduce with initial addition of N. This is the only case known to the authors of alloying actually decreasing disorder.

F. Electrical characterization

The activation energy $\Delta E$ of conductivity of a-C:H:N was derived from an Arrhenius plot of conductivity versus inverse temperature. The Arrhenius plot shows two straight line regimes, one in the temperature range 40–100 °C, and the other between the temperatures of 100–200 °C. The variation of $\Delta E$ shown in Fig. 10 is for temperatures greater than 100 °C. The conduction mechanism for temperatures below 100 °C in a-C:H:N films, where $\Delta E$ varies from 0.34 to 0.55 eV in a similar manner to that shown in Fig. 10,\textsuperscript{20} is attributed to localized tail states. Above 100 °C, thermally activated conduction predominates. For a-C:H:N, which is an amorphous material, the mobilities of electrons and holes at the band edge states are assumed to be similar.

$\Delta E$ for a-C:H (0.5 eV) is considerably less that half the optical gap (1.65 eV). $\Delta E$ then increases to a value of 0.9 eV at 7\% N. This is much closer to half the optical gap at this composition (2.1 eV). $\Delta E$ then declines to 0.46 eV at 15\% N, which is again much less than half the gap (1.8 eV) at that composition. We interpret this as follows. $\Delta E$ corresponds to the energy difference from $E_F$ to the conduction level, usually the nearest band edge. $E_F$ in conventional a-C:H tends to lie near midgap.\textsuperscript{2} However, $E_F$ in ta-C and ta-C:H appears to lie well below midgap, as summarized in the introduction. This arises because half-filled C defect levels lie predominantly below midgap. We believe that the present a-C:H, also with a low H content, has a similar behavior, with its $E_F$ below midgap. In that case, if N is a weak donor, N addition causes $E_F$ to rise through the gap towards $E_C$. This accounts in general terms for the increase and then decrease of $\Delta E$ in Fig. 10. The 7\% N composition is the compensated composition, where the N donors compensate the acceptor-like defects. This situation is locally stable, and perhaps accounts for why the fraction of N sp\textsuperscript{3} bonding reaches a maximum there.

The alternative interpretation, that the activation energy change arises simply from changes in the gap, is rejected because the conductivity changes are consistent with the strong changes in the bonding at N sites shown in Fig. 5.
The compensation occurring at such a high composition as 7% emphasizes the low doping efficiency of N in a-C:H. In essence, the addition of 7% N atoms has moved \( E_F \) through a defect density of say \( 4 \times 10^{18} \) to \( 4 \times 10^{20} \) cm\(^{-3} \) (see Sec. III G), so the efficiency is given by \( 10^{20}/(0.07N_{at}) \) or 0.02%—1.5%, where \( N_{at} \) is the atomic density, \( 3.7 \times 10^{23} \) cm\(^{-3} \) derived from the valence plasmon energy. There are many possibilities for the nature of the N donor level, from Figs. 1(b), 1(c), 1(e), and 1(h).

**G. Electron spin resonance**

Various bonding configurations give rise to states in the gap, which when half-filled and paramagnetic gives rise to an electron spin resonance (ESR) signal. The variation of this spin density as a function of atomic N content is shown in Fig. 11. The spin density decreases rapidly from a high value of \( 4 \times 10^{20} \) cm\(^{-3} \) to values below \( 10^{18} \) cm\(^{-3} \) with increasing N content.

There are two possible reasons for the decrease in spin density, either the total defect density has decreased, or the defects have been converted from their paramagnetic to diamagnetic configuration by the transfer of electrons from N sites. Generally, diamagnetic defects are expected in the presence of dopants, because the donor electrons will fall down from the donor level into the singly occupied defect level and make it fully occupied and diamagnetic. A typical example of this situation is the diamagnetic \( N_3^+ \) configuration shown in Fig. 1(c). This effect has been shown to occur in a-Si:H.\(^{26} \)

On the other hand, the total defect density is correlated with the network disorder and Urbach slope, as occurs in a-Si:H. Thus the decrease in Urbach slope seen in Fig. 9(c) suggests that at least part of the decrease in spin density is due to a decrease in the total defect density. If so, this is a remarkable decrease. We hope to confirm this by measuring the total defect density by light induced ESR, in which all defects are excited to their paramagnetic state.

The nature of the defect configuration can often be identified from the ESR g-factor and linewidth. A g-factor of 2.0028 indicates a carbon center.\(^{39} \) The ESR linewidth in Fig. 11(b), shows it to decrease with increasing nitrogen content. The ESR linewidth in a-C:H films has been attributed to exchange narrowing.\(^{39–43} \) The linewidth under exchange narrowing is proportional to the electron density per site and so inversely proportional to the number of sites over which the unpaired electron is delocalized. The 2.0028 defect has been associated with sp\(^3 \) clusters,\(^{43} \) so the decrease in linewidth with N content suggests that the defect becomes larger with increasing N content. The alternative line broadening mechanism in spin resonance is dipolar broadening by interaction with nearby H atoms, in which the linewidth would increase with H content. A dependence on H content was observed in sputtered a-C:H by Hoinkis et al.\(^{41} \) In our case, the linewidth decreases from 5 to 2 G for N contents from 0% to 7%, while the H content is essentially constant (Fig. 2), which is evidence against this dipolar broadening in PECVD a-C:H.

**IV. DISCUSSION**

The variation of the JDOS as a function of N content is shown in Fig. 12. The area shown boxed in Fig. 12 is expanded in Figs. 12(a) and 12(b). The JDOS for all three films in general can be taken to consist of two broad peaks centered about 5 and 12 eV. The peak at \( \sim 5 \) eV is attributed to \( \pi-\pi^* \) states and the one at 12 eV to \( \sigma-\sigma^* \) states. It is seen that there is a significant reduction in the gap states as the N content changes from 0% to 7%, seen as a reduction in the JDOS in the 0–5 eV region in Fig. 12(a). This is consistent with the widening of the optical gap and sharpening of band tails seen in Fig. 9. As the N content rises to 14%, it is seen that the JDOS at the band edges now increases again. This is attributed to the introduction of extra states near the band edges by N sites.

In analyzing the electronic doping results, a model as to how a predominantly sp\(^3 \) bonded material can contribute to the conduction process must be considered. If it is assumed that each of the N that is sp\(^3 \) bonded to C contributes an electron to the conduction process when found sitting in a
sp\(^3\) C environment Fig. 1(b), and that the N which is sp\(^2\) bonded in a sp\(^2\) C environment contributes an electron due to pairing up of the C \(\pi\) electron with the fourth N electron, Figs. 1(e) and 1(h), then an empirical expression which relates the electronic activation energy for conduction to the C and N bonding configuration can be derived.

\[
\left( k \times N_{\text{sp}^2} \times C_{\text{sp}^3} \right) + \left( m \times N_{\text{sp}^3} \times C_{\text{sp}^3} \right) \times N_{\text{at.}} = \text{constant} \times \Delta E_{\text{intr}},
\]

where \(k\) and \(m\) are the electrically active doping efficiencies in the sp\(^2\) and sp\(^3\) bonding configurations and \(\Delta E_{\text{intr}}\) the movement of the Fermi level from the undoped state. By using a suitable set of axes in the form shown in Fig. 13, \([N_{\text{sp}^2} \times C_{\text{sp}^3} \times N_{\text{at.}} / \Delta E_{\text{intr}}]\) versus \([1 \times N_{\text{sp}^2} \times C_{\text{sp}^3} \times N_{\text{at.}} / \Delta E_{\text{intr}}]\), it is possible to obtain a measure for the relative efficiency of doping \((k/m)\) in the two configurations. A suitable fit to the data shown in Fig. 13 gives a straight line approximation and a value for ‘‘\(k/m\)’’ equivalent to 0.2. The experimental data indicates that it is five times more likely that an N which substitutes in a C–N environment acts as an active donor compared to an N which is in a C–N environment. This result also agrees with the experimental observation that highly tetrahedral a-C can be more easily and efficiently doped with N when compared to a-C:H:N films. In the results published by Veerasamy et al.\(^{14}\) an atomic N content of 0.5% was needed to shift the Fermi level to midgap in highly tetrahedral a-C, whereas an atomic N content of 7% is needed in the a-C:H:N thin films.

The compressive stress in the a-C:H:N films decreased with increasing N content, Fig. 3(b). This variation is evidence for structural relaxation due to the addition of N. The stress in a thin film is a good indicator as to the constraints present in the bonding in a material. Thrope\(^{14}\) defined a mean coordination number \((r)\) for a skeleton network in terms of the fraction of threefold, fourfold coordinated, and H sites in a network,

\[
(r) = \frac{4C_{\text{sp}^3} + 3C_{\text{sp}^2} - C_{\text{H}}}{1 - C_{\text{H}}},
\]

where \(C_{\text{sp}^3}\) is the carbon sp\(^3\) content, \(C_{\text{sp}^2}\) is the carbon sp\(^2\) content and \(C_{\text{H}}\) the hydrogen content in the a-C:H:N films.

Figure 14 shows the variation of the coordination number for the a-C:H:N thin films. The peak in the coordination number is seen to correspond to the maximum of the C and N sp\(^3\) contents. The coordination of our a-C:H:N alloys decreases gradually with N content and rapidly beyond an atomic N content of 7%. This is expected if the atomic N acts as a termination site and thereby relaxes the entire a-C:H:N network towards three. Since the optical band gap still remains relatively high, it is difficult to classify these films as graphitic in nature. As the N bonds preferentially in a threefold coordination in the C–N alloys, the relative N content too should be considered before a classification is made as to the nature of the film.

V. CONCLUSIONS

The effect of nitrogen addition on the structural and electronic properties of hydrogenated amorphous carbon (a-C:H) films has been characterized. The variation of conductivity with nitrogen content suggest that N acts as a weak donor, with the conductivity first decreasing and then increasing as the Fermi level moves up in the band gap. Compensated behavior is found at about 7 at. % N, where various properties such as the optical gap, and N sp\(^3\) content show a maximum. The Urbach energy and paramagnetic defect density are found to decrease with increasing N content. This indicates an unusual decrease in disorder with alloying.


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