Chemically Active Substitutional Nitrogen Impurity in Carbon Nanotubes

Andriy H. Nevidomskyy, Gábor Csányi, and Michael C. Payne
Theory of Condensed Matter Group, Cavendish Laboratory, University of Cambridge, Cambridge CB3 0HE, United Kingdom
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We investigate the nitrogen substitutional impurity in semiconducting zigzag and metallic armchair single-wall carbon nanotubes using \textit{ab initio} density functional theory. At low concentrations (less than 1 at. %), the defect state in a semiconducting tube becomes spatially localized and develops a flat energy level in the band gap. Such a localized state makes the impurity site chemically and electronically active. We find that if two neighboring tubes have their impurities facing one another, an intertube covalent bond forms. This finding opens an intriguing possibility for tunnel junctions, as well as the functionalization of suitably doped carbon nanotubes by selectively forming chemical bonds with ligands at the impurity site. If the intertube bond density is high enough, a highly packed bundle of interlinked single-wall nanotubes can form.

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Ever since their discovery \cite{1,2}, carbon nanotubes have been heralded as the new wonder material of the future. Their remarkable mechanical and electronic properties destine them to play a major role in all kinds of nanotechnologies and molecular electronics. At least two major hurdles have to be overcome in order to fulfill this potential. First, manipulation of individual tubes is at best difficult today, which prevents mass production of devices. Second, the ability to fine-tune the various properties of the material to suit particular applications has to be achieved.

Carbon nanotubes exhibit a variety of electronic properties, for example, depending on their diameter and chirality, they vary from being metallic to semiconducting \cite{3}. The presence of defects and impurities that are electronically or chemically active can change these properties and thus have a significant bearing on a broad range of applications. In this paper we investigate the nitrogen substitutional impurity in single-wall carbon nanotubes (SWNT). In the case of semiconducting tubes, doping with electrons or holes is the principle route to making electronic devices. Alternatively, introducing new levels in the band gap with associated electronic states that are spatially localized can create chemically active impurity sites.

Several groups have produced nanotubes containing nitrogen \cite{4–14} or nitrogen and boron \cite{5,15,16}. All of these were multiwall tubes, except for \cite{16}, where nitrogen was always accompanied by boron. The measured atomic concentrations of the impurities were relatively high in all cases, above 1%, and all the CN\textsubscript{x} tubes were found to be metallic, as predicted by theoretical models \cite{10,11,17}. Indeed, one other reason for the particular interest in CN\textsubscript{x} tubes was that they might make even initially semiconducting tubes metallic.

Various structural models have been proposed for the incorporation of nitrogen into the carbon network. Direct substitution was first studied by Yi and Bernholc \cite{18} using density functional theory (DFT) at an atomic impurity concentration of about 1%, where they found an impurity state lying 0.27 eV below the bottom of the conduction band at the $\Gamma$ point and a significant overlap of the states associated with the adjacent nitrogens (8.4 Å apart). Lammert \textit{et al.} \cite{19} considered disordered substitution of boron and nitrogen at a concentration of 3.5% and 1.4%, and observed both donor and acceptor levels in the gap. For an isolated nitrogen or boron impurity, they found the impurity state to be localized within about 10 Å for an (8,0) tube.

Another atomic arrangement for the experimentally observed nitrogen impurity was proposed by Terrones \textit{et al.} \cite{6}, where they suggest that at high concentrations (above about 10%) nitrogens are divalent, pyridinelike. Later publications from the same group refine this model and correlate the calculated density of states of metallic tubes with experimental observations \cite{10,11}.

To study the isolated substitutional nitrogen defect, the simulation cell has to be as long as possible. The existing constraints of computational power mean that the diameter of the tube should therefore be as small as possible. In all calculations we used the achiral zigzag (8,0) and armchair (5,5) nanotubes. Our simulation cell has periodic boundary conditions; the tubes are arranged parallel to the $z$ axis and form a triangular lattice in the perpendicular $x$-$y$ plane. This arrangement is experimentally seen in carbon nanotube ropes \cite{3}. We studied a range of nitrogen concentrations from 0.26% to 1% by introducing one impurity atom and changing the size of the simulation cell along the axis of the tube. The largest simulation cell contained 12 unit cells along the $z$ axis for the zigzag tube and 14 unit cells for the armchair tube, respectively, with one impurity atom in it.

Throughout, we used \textit{ab initio} plane-wave pseudopotential density functional theory \cite{20} as implemented in the \textit{CASTEP} code \cite{21}. The generalized gradient approximation was used to account for exchange and correlation in the Perdew-Burke-Ernzerhof form \cite{22}. All calculations were spin polarized and ultrasoft
pseudopotentials were used for carbon and nitrogen with a cutoff energy of 300 eV. We assessed the adequacy of this cutoff by performing calculations at 350 eV for the smallest unit cells and verified that none of the properties reported in this work changed perceptibly. For each concentration, the ionic positions were optimized by using one $k$ point at the $\Gamma$ point for the largest simulation cells and correspondingly more $k$ points for the smaller cells. After the geometry optimization converged, the band structure was calculated using a finer mesh of $k$ points.

We find that the equilibrium position of the nitrogen ion is almost unchanged with respect to the corresponding C atom in the undoped nanotube, being moved by at most 0.01 Å, similarly to the case of azafullerene, C$_{59}$N [23,24]. A slice of the calculated spin density for both zigzag and armchair cases is shown in Fig. 1. The spin density is maximal on the N ion and the neighboring C ions and shows oscillatory behavior as it reaches a local maximum on every other ion in the network, again similarly to that observed in C$_{59}$N [23,24]. The shape of the spin density isosurfaces is reminiscent of the $\pi$ orbitals in graphite with their axes of symmetry perpendicular to the sheet.

The localization of the extra electron associated with the isolated impurity shows a striking difference between the metallic and semiconducting cases, despite the fact that the band gap of the latter is quite small (0.6 eV according to the DFT calculations). In the armchair tube, the unpaired electron is almost completely delocalized along the tube, as can be seen from the plot of the envelope of the spin density in the lower panel of Fig. 1. However, for the zigzag tube, it is localized near the impurity site, but the localization region is fairly large, about 30 Å. This region contains about 90% of the unpaired spin density. The decaying envelope of the spin density is exponential.

The oscillations in the spin density can be understood as follows. Regarding the nitrogen atom as a perturbation, the response of the system is characterized by the static susceptibility $\chi(r)$. The characteristic period in $\chi(r)$ corresponds to a maximum in the Fourier transform, $\chi(q)$, which happens for $q$ values that connect points in the Brillouin zone with the minimum energy difference. For the 2D graphene lattice, the energy gap is zero at the high symmetry $K$ points, so for this and similar systems, $\chi(r)$ will oscillate with a period corresponding to the momentum transfer between different $K$ points, which is what is observed. The well-known oscillations in the susceptibility in the RKKY formalism [25] are essentially a special case of the above considerations.

The decay of the envelope of the spin density is also analogous to the RKKY case. For metals in 1D, the result is $\chi(r) \sim 1/r$ [26,27]. For semiconductors, an analogous coupling to excited states takes place, and assuming a parabolic conduction band, it was shown [28] that $\chi(r) \sim \exp[-(\alpha E_{\text{gap}})^2 r]$, where $\alpha = (2m_{\text{eff}})^{1/2}/\hbar$. For the zigzag nanotube, using the calculated value of $m_{\text{eff}}/m_e = 0.2$ this crude estimate gives a decay constant of about 6 Å, close to the observed 7 Å.

Note that in all the graphene-based systems (C$_{59}$N, nitrogen-doped graphene and nanotubes), the spin density is positive almost everywhere. At first sight, this might seem to contradict the results from electron spin resonance hyperfine measurements, which indicate that the spin density oscillates symmetrically around zero. In fact, as was mentioned above, the spin density is very strongly dominated by the $p$ component, whereas the Fermi contact interaction in the hyperfine coupling is sensitive only to the $s$ component, so its observed symmetrical oscillation is consistent with the calculations.

A crucial difference between the armchair and the zigzag tubes is that the introduction of the nitrogen atom breaks the left-right mirror symmetry in the latter case. As two impurity atoms approach each other in an armchair tube, their interaction will show oscillatory behavior depending on whether the spin density maxima induced by the respective nitrogens coincide or not. But in the zigzag tube the local maxima of the spin density occur on different sublattices on the two sides of the nitrogen atom. As two impurities approach, their respective spin density maxima never coincide, irrespective of whether the nitrogens occupy the same sublattice or not.

Figure 2 shows the band structure of the nitrogen-doped nanotubes along with the corresponding densities of states. The band structure of the undoped tube (folded in such a way that it fits into the shrunk Brillouin zone corresponding to the larger supercell) is shown for

![Figure 1](color). Slice of the spin density of the N-doped carbon nanotube in a plane parallel to the tube axis, the plane being tangential to the tube at the position of the nitrogen ion, for (a) (8,0) zigzag tube, comprising 12 unit cells (50.6 Å), and (b) (5,5) armchair tube, comprising 14 unit cells (34.3 Å). Above each slice, we plot the envelope of the spin density (integrated in the plane perpendicular to the tube) in the units of electron/Å. The nitrogen ion is marked white. The color range maps the value of the spin density, with blue corresponding to the smallest and red to the highest values. Note that the scale of the spin density is quite different for the two panels.
comparison. For the zigzag case, the extra band of the impurity falls somewhere near the bottom of the conduction band, and a complicated hybridization takes place with the existing unoccupied bands. The end result is that a new flat level appears about 0.2 eV below the bottom of the conduction band. In the armchair tube, the impurity introduces a peak at about 0.6 eV below the first van Hove singularity above the Fermi level, as well as a number of additional peaks, which are formed because of the hybridization and level repulsion taking place between the energy levels of impurity and pristine nanotube.

The localized electronic state of the semiconducting tube offers the intriguing possibility of being chemically active and forming covalent bonds with ligands or other tubes. Figure 3 shows two zigzag nanotubes each doped with nitrogen (one per four unit cells) in such a way that two carbon atoms with originally significant spin density on them face each other. As the charge density slice shows, a covalent bond is formed between these atoms, and thus between the tubes. The two carbon atoms pop out of the tangent plane of the tubes, and take up an \( sp^3 \) configuration, the bond length between them being 1.65 Å (note that a similar geometry was discussed in the context of carbon nitride materials [29]). In order to obtain the geometry shown, the two tubes were pressed against each other, which is realistic when modeling a nanotube lying on top of another on a surface [30]. The further corroboration of the formation of a true covalent bond between the carbon atoms comes from the analysis of the charge density originating from different orbitals. The largest contribution to the total charge density between the newly bonded carbon atoms comes from the highest occupied molecular orbital (HOMO) [Fig. 3(b)], which is occupied by about 0.3 of an electron.

On the other hand, for the intertube bonds to be realized in a bundle of SWNTs, they have to be stable under zero external pressure. We found that this is possible if the number of intertube bonds is increased to one per two unit cells (8.4 Å), so that enough energy is gained in the new bonds to overcome the intertube repulsion. Here, the bonded configuration is a local minimum and the tube–tube distance is reduced to 2.5 Å from 3.4 Å in the case of the unbonded bundle. This finding opens the possibility of producing highly packed cross-linked bundles of doped nanotubes; we obtained a crude estimate of 80 GPa for the bundle shear modulus (as a result of just one interlink for every two unit cells), which is substantially larger than 1–6 GPa, reported for ordinary bundles [31].

Figure 4(a) shows the \( p \)-projected local density of states (DOS) of various carbon atoms in different sections of the nanotube with one intertube bond per two unit cells, as compared to that of the unbonded system. The dashed curve shows the fine structure of the \( \pi^* \) peak,
which is almost completely absent for the $sp^3$ carbon atom (solid curve) that takes part in the intertube bond. As a first approximation, the $p$ projected total DOS in the conduction band, when suitably shifted, is proportional to the electron energy-loss spectrum (EELS) [32,33], shown in Fig. 4(b). In the bonded system the $\pi^*$ peak is somewhat diminished, but more importantly, the peak marked with an arrow is absent. Since the proportion of $sp^3$ bonded carbon atoms is expected to be quite small, a local probe such as EELS would be most suitable for the detection of the intertube bond.

We performed ab initio calculations of zigzag and armchair carbon nanotubes substitutionally doped with nitrogen. In the case of the armchair nanotube, the impurity state is totally delocalized and the corresponding energy level falls inside the unoccupied bands of the pristine nanotube. In the case of the semiconducting zigzag nanotube, however, the isolated nitrogen impurity forms a flat energy level lying inside the band gap, 0.2 eV below the bottom of the conduction band. It hybridizes with the $\pi$ orbitals to create a spatially localized state decaying exponentially with an overall extent of about 30 Å. This singly occupied state is chemically active, and we demonstrate the possibility of the formation of a covalent bond between two nanotubes with impurities that face one another. This opens the intriguing general possibility for chemical bonding between suitably doped carbon nanotubes or indeed with other ligands. Such tube-tube junctions could be interesting from a device perspective as the intertube bond will probably change the tunneling properties between the tubes considerably. If the density of intertube bonds is high enough, a bundle of SWNTs can become highly interlinked and packed, substantially enhancing its mechanical properties. The ligand-binding properties of the localized impurity site, of interest in chemical sensing applications, are currently under investigation.

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