Metallic behaviour of carrier-polarized C\textsubscript{60} molecular layers: Experiment and Theory

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Although C\textsubscript{60} is a molecular crystal with a bandgap $E_g$ of $\sim 2.5$ eV, we show that $E_g$ is strongly affected by injected charge. In sharp contrast to the Coulomb blockade typical of quantum dots, $E_g$ is reduced by the Coulomb effects. The conductance of a thin C\textsubscript{60} layer sandwiched between metal (Al, Ag, Au, Mg and Pt) contacts is investigated. Excellent Ohmic conductance is observed for Al electrodes protected with ultra-thin LiF layers. First-principles calculations, Hubbard models etc., show that the energy gap of C\textsubscript{60} is dramatically reduced when electrons hop from C\textsubscript{60} to C\textsubscript{60}.

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The Fullerene C\textsubscript{60} solid is a molecular crystal with a bandgap of $\sim 2.5$ eV. Alkali-metal doping makes it a conductor or a superconductor. Other methods of converting C\textsubscript{60} into a conductor involve exotic chemical routes where, e.g., C\textsubscript{60} molecules are joined by metal-ligand structures. Such efforts focus on improving the conductance of the molecular layer itself. Another important factor is the electrical contact between the molecular layer and the metal surface. The practical realization of molecular electronics depends crucially on this molecule/metal contact. In many situations, such contacts involve chemical processes that lead to fragmentation of the molecules themselves.

Fullerenes, e.g., C\textsubscript{60} are popular candidates for molecular electronics. Unlike quantum dots and related nanostructures, molecules come in identical copies and have energy levels which are robust at room temperature. Many studies on molecular-C\textsubscript{60} devices have emphasized negative-differential resistance (NDR), or the control of conductance using scanning-tunnelling probes, electromechanical and other gates. In the simulation by Taylor et al., a single C\textsubscript{60} molecule is positioned between two Al leads. In their ideal device, metallic conductance occurs when the triad of degenerate LUMO (lowest-unoccupied molecular orbital) acquires three electrons and catalyse the decom-

Studies of C\textsubscript{60} layers deposited on crystallographically controlled surfaces of Al, Ag, Au, Mg, etc., show that charge transfer, electronic and bonding modifications occur within the first monolayer of molecules, but the second monolayer usually remains unaffected, except for weaker physical effects. More than one C\textsubscript{60} monolayer may be affected if the surface is polycrystalline. We call this physico-chemically modified interface-layer the metal-fulleride layer. The metal-fulleride formation for C\textsubscript{60} deposition on Al, as well as for Al deposition on C\textsubscript{60} has been studied by, e.g., Owens et al. The interaction is more complex than resulting from image-charge effects or simple charge transfer, as may perhaps be the case for Ag. The situation is even more complicated for Pt and Ni surfaces which form strong bonds with C\textsubscript{60} and catalyse the decomposition of the organic molecule at sufficiently elevated temperatures. If the “bulk” C\textsubscript{60} material is clean and undoped, very few carriers are available for producing “band-bending” effects of dipole layers etc., as found at doped inorganic semi-conductor interfaces. This implies that, except for the fulleride layer next to the electrode, the molecules in the “bulk” are unaffected by the presence of the metal contacts. This picture assumes that hot-metal atoms were not bombarded into the organic film, or that the metal was not presented as a paste or other preparation where metal diffusion into the C\textsubscript{60} could occur.

Most reports of I–V behaviour on such metal/C\textsubscript{60}/metal sandwich structures suggest semiconducting, rectifying or insulating behaviour. In a previous paper, we showed how an ultra-thin layer of LiF dramatically modifies the I–V characteristics of a molecular film by (i) protecting the organic molecules from reactions with the metal, (ii) creating a sharp electrode density of states (DOS) and favouring carrier injection. Similar important effects of ionic epilayers on metals are not

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uncommon\cite{18}. Here we study C$_{60}$ molecular films and show that the charge injection dramatically changes the “insulating” or semi-conducting character of C$_{60}$ films. Metallic conductance is achieved for M = Al and Mg. That is, the current $I$ depends linearly on the applied bias $V$, for both forward and reverse bias. When M=Pt, the conductance is very small, non-linear and there is strong asymmetry on reversing the bias. We note that the work functions for polycrystalline Mg, Ag, Al, Au and Pt are 3.7, 4.3, 4.3, 5.1 and 5.7 eV respectively\cite{19}. Although there is some correlation with the work function, the situation is more complex since carrier hopping is subject to Coulomb interactions as well.

Even if the C$_{60}$ monolayer (fulleride layer) just next to the metal electrode acquires electrons due to interactions with the metal\cite{11}, conduction cannot occur unless there is a mechanism for further charge transfer from molecule to molecule\cite{20} in undoped C$_{60}$. We clarify the observed metallic behaviour using detailed electronic structure calculations. The energy gap of the “bulk” C$_{60}$ molecules dynamically collapses as the carrier electrons hop from molecule to molecule. That is, an inverse coulomb blockade occurs, where charge transfer between molecules is enhanced. In metals like Pt where fulleride formation via metal d-orbitals is likely, charge injection is negligible under low bias; here strongly non-linear $I$–$V$ is observed. Thus while neutral C$_{60}$ behaves as a molecular crystal in photoemission experiments\cite{1}, our experiments and calculations suggest that C$_{60}$ may behave as a metal in the presence of polarizing carriers.

**Experimental.**—Details of sample fabrication etc., are given in Lu et al.\cite{17}. The devices were made on 2 in x 2 in Si(100) wafers with 2000 nm furnace oxides on top. The first metal (Al, Mg, Ag, Au, Pt) electrode (1 mm wide, 50 mm long, and 60 nm thick) was deposited through a shadow mask, and is referred to as the “bottom” electrode. C$_{60}$ films (210 nm thick) were then deposited over the bottom electrode. A second metal (Al) electrode (1 mm wide, 50mm long and 100 nm thick), referred to as the “top electrode”, was deposited over the C$_{60}$ films. An ultra-thin layer (~ 1 nm) of LiF was deposited on the C$_{60}$ film prior to the deposition of the top electrode. The top electrode lines were orthogonal to the bottom electrode lines so that each intersection of these two lines produces one metal/C$_{60}$/metal device. There are 20 to 100 devices on each wafer. A final silicon oxide film of ~ 300 nm was used to encapsulate the device. This final encapsulation is essential to produce consistent and reproducible results. Such protection is important since contamination with air has a drastic effect\cite{14} on the electrical properties of C$_{60}$. All devices were made using a K.J. Lesker 4 in x 4 in Cluster Tool having several process chambers inter-linked through a central distribution chamber. All metals were deposited in the metallization chamber having a base pressure of ~10$^{-7}$ Torr. The C$_{60}$ molecules were deposited in the organic chamber having a base pressure of ~ 10$^{-8}$ Torr. The transfer of samples between chambers was via a central chamber, with a base pressure of ~ 10$^{-9}$ Torr. The $I$–$V$ measurements were in a dark, ambient environment, using an HP4140B meter with a Materials Development probe station.

Figure 1, top panel shows the $I$–$V$ data for samples which differ in the presence or absence of LiF interlayers. In samples with bare-Al electrodes, i.e., Al/C$_{60}$/Al the left-hand Al (the “top” electrode) was deposited hot on C$_{60}$, while the right-hand Al electrode (“bottom” electrode) was cold during C$_{60}$ deposition. In such systems the fulleride interlayers are uncontrolled and they are denoted by //, a double bar. In devices with LiF at the top electrode, i.e, Al/LiF/C$_{60}$/Al, the C$_{60}$ is protected from the hot deposition of the “top” Al electrode.

The device with bare Al electrodes produces a typical diode-like $I$–$V$ behaviour (see Fig. 1, top panel) with negligible current flow at low bias. The reverse-bias current flow “takes off” more rapidly, as seen in the figure. The use of a LiF interlayer at the top electrode (the “hot-deposited” electrode) is enough to create a linear $I$–$V$ relationship with zero threshold for current flow. This is typical of metallic conductors. When a LiF interlayer is included at the bottom electrode as well, the conductance ($I/V$) improves only slightly, showing that the bottom electrode, where C$_{60}$ was deposited in the cold, required no LiF protection. The behaviour is completely metallic, with no discontinuity in the gradient $I/V$ on changing the sign of $V$. This was confirmed for C$_{60}$ layer vary-
ing from 100 nm to 250 nm in thickness. It is clear that Al$^{3+}$ migration into the C$_{60}$ layer cannot be the cause of the conductance (as was the case in Ref. [12]), since such migration is obstructed by the LiF layer. Photoemission experiments suggest that only the first few monolayers of C$_{60}$ near the metal are affected by the presence of the metal[8]. The $I - V$ data show that fulleride formation at the cold-deposited C$_{60}$ on the Al interface is not critical to electron transport, while the prevention of fulleride formation at the hot-deposited Al on C$_{60}$ is absolutely important. The improved conductance of the sample with the Al electrodes protected with LiF confirms the protective role of LiF. This also suggests that the resistance of the device is determined by that of the fulleride layer which plays the role of the least conducting link.

Figure 1, bottom panel shows typical results for Al, Ag and Pt. Both Al and Mg (not shown) show metallic behaviour. However, surprisingly, Ag shows some slight non-linearity in its $I - V$, revealing some interface reactivity[9]. Results for Au (not shown) tend to be significantly more non-linear than for Ag. The Pt/C$_{60}$ interface is clearly very unfavourable for Ohmic $I - V$.

Theory.– We have studied the HOMO-LUMO energy gap $E_g$ as well as the electronic density of states (DOS) for the C$_{60}$ molecule, for C$_{60}$-LiF, C$_{60}$-C$_{60}$, C$_{60}$-C$_{60}$ and C$_{60}$-C$_{60}$ structures, where the short bar stands for all the interactions between the two moieties. These electronic-structure details are obtained from density functional calculations using the Gaussian-98 code[21] at the BP86/6-31G* level.[21, 22]. The calculations included geometry optimization via total energy minimization using gradient corrected exchange-correlation functionals. The modifications in the DOS of some of the C$_{60}$ structures are shown in Fig. 2 and in Table I. In the upper panel we compare our isolated-molecule DOS with the experimental results[6] for multilayers of C$_{60}$ on Ag obtained using direct and inverse photoemission (UPS and IPS). Thus the HOMO part of the experimental curve is from UPS (energy resolution $\sim 0.1$ eV), and the LUMO part of the curve is from IPS (energy resolution $\sim 0.3$ eV). We have positioned these experimental curves so that our calculated HOMO (H) and LUMO (L) peaks align with the experimental H and L peaks. The general agreement clearly confirms that the molecules in a C$_{60}$ multilayer deposited on Ag are essentially like isolated molecules. This also serves to confirm the validity of our theoretical calculations.

The presence of LiF itself produces a decrease in the C$_{60}$ energy gap. Here the distance between the LiF and the C$_{60}$ center, as well as other inter-nuclear distances have been energy optimized, and hence this is the upper-bound to the gap reduction that may arise from LiF.

The energetics of the C$_{60}$ interactions is given in Table I. The energy gaps $E_g$ given in the table improve on the gaps calculated using the local-density approxim-

| system | $E_H$ a.u. | $E_L$ a.u. | $E_g$ eV |
|--------|-----------|-----------|--------|
| C$_{60}$ | -0.22003  | -0.11854  | 2.8    |
| C$_{60}$-LiF | -0.18993 | -0.16157  | 0.8    |
| C$_{60}$-C$_{60}^\alpha$ | -0.11880 | -0.06443  | 1.48   |
| C$_{60}$-C$_{60}^\beta$ | -0.11736 | -0.06213  | 1.50   |
| C$_{60}$-C$_{60}^\alpha$ | -0.28683 | -0.23093  | 1.52   |
| C$_{60}$-C$_{60}^\beta$ | -0.28485 | -0.22942  | 1.51   |

mation (LDA). Thus the LDA gap for C$_{60}$ is only $\sim 1.8$ eV, and differs from the experimental solid-state gap of $\sim 2.5$ eV. Our spin density-functional calculations show a spin-polarized energy-level structure for the anion. The electrons of the C$_{60}$ system form a low-density quasi-2D electron fluid. Such systems have spin dependent ground states under suitable conditions[23]. This may be of importance in spin transport applications using suitable C$_{60}$ based systems[24], although here the spin splitting is only about $\sim 0.02$ eV.

Photoemission experiments[8, 9] clearly establish the metalization of the first C$_{60}$ layer on metals like Ag and Al, with as much as 1.8 electrons transferred[27] to the LUMO in the case of Ag. That this is close to 2 electrons per molecule suggests that the Hubbard like on-site repulsion has been reduced from that of the isolated molecule. An electron transferred to a C$_{60}$ molecule, and subject to the applied bias, hops to an adjacent C$_{60}$ when carrier transport occurs. The final state of such a hop is given in our calculations for the (C$_{60}$-C$_{60}$)$^-$ anion, and shows a reduced gap. This is due to (a) splitting of the five-fold HOMO and the three-fold LUMO multiplets by the electric field of the in-coming charge, (b) polarization and distortion of the of the molecules which persist under stationary state conditions, (c) resulting modification of the on-site Coulomb interactions. Many of these issues have been examined using Hubbard type models[20], or with microscopic approaches. Hesper et al. [20] used an image-charge model for C$_{60}$ on Ag to obtain a gap reduction of $\sim 1.44$ eV. They even mention the possibility of "driving the insulator into the metallic state". Our explicit calculations (Table I) can be used to parametrize the Hubbard models. However, even the Hubbard-model conductance has not been evaluated, except in special cases (e.g. infinite-dimensional Hubbard models).

In the usual picture of electron transport across molec-
A transient, excited electron is injected from the source electrode to the LUMO of the nearest molecule, converting it to an electron that an electron is injected from the source electrode to the LUMO of the nearest molecule, converting it to an electron. The usual factor, viz., the transmission coefficient $T_{fb}$ for the process, metal-fulleride $\rightarrow C_{60}$ bulk-like layer. An evaluation of $T_{fb}$ is postponed till a detailed fulleride $C_{60}$ calculation becomes available.

Our geometry optimizied calculations (Table I) includes the electron-phonon interaction to all orders. The importance of these effects is recognized, especially within the alkali-doped fullerides like $K_{x}C_{60}$ which is an insulator due to bond-distortion effects. Also, $K_{6}C_{60}$ is a band insulator since the 6 electrons per molecule completely fill the three-fold LUMO. In our system, the LUMO of the fulleride layer is only occupied to $\leq 1.8$ electrons per $C_{60}$ molecule.

In conclusion, we have shown, experimentally and theoretically, that $I - V$ characteristics similar to a metal can be obtained using LiF protected low-workfunction electrodes connected to $C_{60}$ layers.

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