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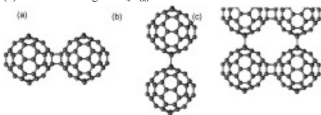
# Mixed Interfullerene Bonding Motifs in $C_{60}$ based polymers

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An unexpected discovery in fullerene chemistry has been the ease with which C<sub>60</sub> units can covalently bond together to give rise to polymerized fullerene networks with a variety of structural architectures. Such fullerene-bridged arrays display varying dimensionality and interesting electronic (metallic behavior) and magnetic (ferromagnetism above room temperature) properties. The predominant C-C bridging structural motif, encountered in photo- and pressure-polymerized neutral C<sub>60</sub> and in the  $AC_{60}$  (A = K, Rb, Cs) solids arises from [2+2] cycloaddition reactions, which result in the formation of 4-membered carbon rings, fusing together adjacent molecules and propagating in one (1D chains) or two (2D layers) dimensions (Fig. 1a). An alternative bridging mechanism involves the formation of single interfullerene C-C covalent bonds, as encountered in 1D  $C_{60}^{3-}$  (Na<sub>2</sub>RbC<sub>60</sub>) [1] (Fig. 1b) and 2D  $C_{60}^{4-}$  (Na<sub>4</sub>C<sub>60</sub>)<sup>3</sup> [2] fulleride polymers. We have recently started working on the related fulleride salt, Li<sub>4</sub>C<sub>60</sub> and its structural properties were probed by X-ray powder diffraction as a function of temperature between ambient conditions and 450(C. Very surprisingly we found that at room temperature the ground state of Li<sub>4</sub>C<sub>60</sub> is a two-dimensional polymer with monoclinic crystal symmetry and an unprecedented architecture, combining both [2+2] cycloaddition and single C-C bridging motifs [3,4] (Fig. 1c). This structure is the first example of a fullerene polymer with a mixed mode of interfullerene bridging and opens the way for the synthesis and study of new  $C_{60}$ -based 1D, 2D and 3D polymers. Increasing the temperature leads to the disruption of the bridging motifs and to the simultaneous formation of a fcc monomeric phase.

Figure 1. Schematic drawing of the interfullerene C-C bridging structural motifs in polymeric fullerides. (a) [2+2] cycloaddition in  $AC_{60}$ , (b) single C-C covalent bonds in  $Na_2RbC_{60}$ , and (c) mixed bonding in  $Li_4C_{60}$ 



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# Nanotube field of $C_{60}$ and $C_{70}$ molecules in carbon nanotubes

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We present calculations of the interaction energy of  $C_{60}$  fullerene molecules encapsulated in single-walled carbon nanotubes (SWCNTs), systems known as peapods [1]. We focus on the dependence on the molecular orientations by taking the molecular structure fully into account. First, we approximate the SWCNT as a homogenous cylindrical distribution of carbon atoms, hence keeping the tube's radius R as a single tube-characteristic parameter rather than the two chiral indices n and m [2]. For molecules aligned on the tube axis, we find three different preferential molecular orientations for tube radii in the range 6.5 Å < R < 8.5 Å. We also consider the possibility of having off-centre molecular positions and provide a theoretical description of  $(C_{60})_n@SW\hat{C}NT$  peapods as one-dimensional systems. Then, we test our homogenous-tube approximation by calculating the same  $C_{60}$ -tube interaction for several actual SWCNTs having tube radii in the same range and investigate its validity [3]. We apply the same approximation to  $(C_{70})_n$ @SWCNT peapods. Again, for molecules positioned centrally in the tube, we obtain radius-dependent preferential orientations. "Lying" orientations (long molecular axis coinciding with the tube axis) occur at the lower radii, higher radii yield "standing" orientations (long molecular axis perpendicular to the tube axis), consistent with experimental electron diffraction results [4].

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