of the apex BH unit (decapitation) in these species generates the respective open-faced complexes Cp*CoRR'C2B3H5 (6-8) whose conversion to larger systems (e.g., multidecker sandwiches) is anticipated via deprotonation and reaction with transition-metal ions.2-4

This chemistry, in combination with the previously reported regiospecific B-functionalization of LM(R₂C₂B₃H₅) complexes, ^{2c,d} allows the introduction of desired substituents at specific boron and/or carbon locations and thereby considerably augments the utility of such species in constructing large multimetal systems. Such derivatization at boron has already been exploited to create B-X-B linked oligomers and to electronically tailor the metalcomplexing properties of the open C₂B₃ rim, e.g., in the recent synthesis4 of tetradecker sandwiches via introduction of electron-withdrawing substituents at boron. In addition to the synthetic advantages, the electronic consequences of placing such groups at the cage carbon locations are clearly of interest and are among the numerous ramifications of these findings that we are currently exploring.

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Department of Chemistry University of Virginia Charlottesville, Virginia 22901 Mark A. Benvenuto Russell N. Grimes*

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First Easily Reproduced Solution-Phase Synthesis and Confirmation of Superconductivity in the Fullerene K_xC₆₀ $(T_c = 18.0 \pm 0.1 \text{ K})$

The number of novel molecular superconductors has risen markedly in recent years, especially in the case of the radical-cation ET-based organic materials [ET is the abbreviation for BEDT-TTF, bis(ethylenedithio)tetrathiafulvalene]. The highest confirmed superconducting transition temperatures (T_c) for the organic materials occur for the salts κ -(ET)₂Cu[N(CN)₂]X, X = Br $(T_c = 11.6 \text{ K})^2$ and X = Cl $(0.3 \text{ kbar}, 12.8 \text{ K}).^3$ Very recent work has now established an even higher T_c for a molecular superconductor in the anion-based fullerene (buckminsterfullerene, C_{60}) system $K_x C_{60}$, with onset $T_c \approx 18 \text{ K.}^4$ Because many higher fullerenes are known to exist,5 this new system provides a fertile field for future research on novel superconducting materials. In this communication we report the first easily reproduced solution-phase synthesis of K_xC_{60} and the confirmation of T_c at 18.0 ± 0.1 K. It is especially noteworthy that we find that reduction of C60 in solution with excess K does not lead to a highly resistive material but rather to superconducting K_xC_{60} , in direct contrast to the K-vapor synthesis originally reported.4

Synthesis of C₆₀. Soot containing C₆₀ was prepared in an apparatus described previously.6 Our method is similar to the contact arc method reported by Haufler et al.7 but uses a plasma

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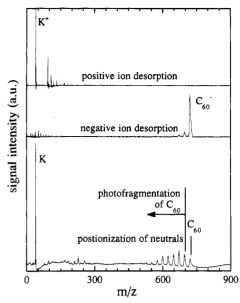


Figure 1. Laser-desorption TOF mass spectra of the K_xC₆₀ sample from a stainless steel substrate with 50-ps 266-nm laser radiation. The top panel shows positive ion desorption, the center panel negative ion desorption, and the bottom panel postionization of desorbed neutrals with 308-nm 10-ns laser radiation.

instead of a contact arc to generate the soot. Current (80 A at 20 V) was passed through the graphite (National Carbon Co., spectroscopic grade) electrodes and ignited a plasma. The plasma was observed through a viewport, and the gap between the electrodes was continuously adjusted to attain maximum plasma brightness. Fine control of the gap distance resulted in a higher yield (up to 44%) of soluble material in the soot.⁶ The soot was collected on a shim cooled by contact with a water-cooled shield. After each run, the shim was removed and the soot scraped from it and dissolved in toluene. Pure C₆₀ was obtained from this mixture by chromatography on a neutral alumina column, as reported previously.8 Laser desorption time-of-flight mass spectrometry9 of the C60 starting material showed that only C60 was present. No C₇₀ impurity was detected at a level above the detection limit of 0.2%.

Synthesis of K_xC₆₀. The preparation was carried out with a Schlenk apparatus under airless conditions. A 40-mL toluene solution containing ~ 15 mg of pure C_{60} (0.021 mmol) was freeze-thaw-degassed three times. Small potassium chips (~75 mg, 1.92 mmol, 90-fold excess) were added while the C₆₀/toluene solution was kept frozen. The reaction flask was immediately evacuated and back-filled with Ar three times. The mixture was warmed to room temperature and then refluxed for 2 h with vigorous magnetic bar stirring. The solution color turned from purple (pure C₆₀) to burgundy and finally to black with a large amount of black precipitate being formed. It was filtered to remove the almost colorless toluene solution and vacuum-dried at room temperature. The Schlenk flask containing the dried black powder was transferred to an Ar-filled drybox, and any large potassium beads were separated. The black powder (K_xC₆₀) containing tiny (nonsuperconducting) potassium beads was loaded into various sample containers and sealed under either Ar or vacuum for further physical characterization (vide infra). The same material could be formed also with a 180-fold excess of K.

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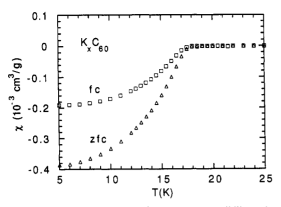


Figure 2. Temperature dependence of the mass susceptibility χ (cm³/g) at an applied magnetic field of 10 Oe for a field-cooled (fc) and zerofield-cooled (zfc) sample of K_xC_{60} , showing T_c onset = 18.0 ± 0.1 K.

Mass Spectral Analysis of K_xC_{60} . The K_xC_{60} compound was placed on a stainless steel sample mount and inserted into a laser desorption time-of-flight mass spectrometer. The top two panels of Figure 1 display prompt ion desorption utilizing 266-nm, ≈50-ps laser pulses (positive ions on top, negative ions on center panel). The bottom panel displays postionization of the laser-desorbed neutrals using 308-nm, ≈10-ns laser pulses. In the positive ion spectrum the prominent peak is K+ with some impurities from the sample mount. No C₆₀⁺ was seen, which is in contrast to the positive ion spectrum from the original C₆₀ sample. This implies that the precursor state for direct ion desorption of C₆₀⁺ was different for the pure and the doped sample. This further suggests that K_xC₆₀ has an ionlike bonding, with C₆₀ forming the anion and K the cation. In the negative ion spectrum we observe only C₆₀, with little fragmentation showing up on the low-mass side as C_n clusters (n = 1-10). Finally, the postionized spectrum shows C₆₀⁺ and K⁺, but, unfortunately, no compound molecule of the kind K_xC_{60} . C_{60} exhibits the usual photofragmentation behavior that has been studied previously.¹⁰ Since no K_xC_{60} molecule is observed directly in the mass spectra, it is not possible at present to quantify the stoichiometry of the solid, but some evidence for the existence of a K_xC₆₀ molecule is given by the three mass spectra. Furthermore, the postionization mass spectrum hints that some of these molecules may laser-desorb intact, but postionization leads to detachment of (all) the potassium(s) even before the C₆₀ molecule starts to fragment. This suggests that the potassium is less strongly bound to the C₆₀ molecule, compared to the bonding of carbon atoms within the C₆₀ molecule. Finally, we see no evidence for C₆₀H_n species, at the 1% level, that might be expected from a modified Birch reduction.

Superconductivity. Superconductivity of the bulk K_xC₆₀ sample was confirmed by both rf penetration depth measurements 11,12 and low-field dc magnetization measurements. Both methods gave comparable results of $T_c \approx 18$ K, but the latter method provides more quantitative information. The magnetization measurements were carried out with the use of a superconducting quantum interference device (SQUID) magnetometer. For these measurements, the black polycrystalline K_xC₆₀ powder was sealed under vacuum in a quartz ampule, and the magnetization was determined for both zero field (zfc) and field cooling (fc) in a magnetic field of 10 Oe. The magnetization curves, expressed as the mass susceptibility $\chi = M/HW$, where M is the magnetization, H the magnetic field, and W the mass of the sample, are illustrated in Figure 2. This figure shows a diamagnetic onset

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of 18.0 ± 0.1 K. The curves are very comparable to those reported by Hebard et al. The shielding (i.e., zero-field cooled) susceptibility at 5 K corresponds approximately to 1% of the value for a perfect superconductor $(-1/4\pi)$, on the assumption of a density of ~2 g/cm³. This indicates that only about 1% by volume of the specimen is superconducting, inasmuch as the shielding volume susceptibility (zfc) of a pure superconducting phase is expected to be close to 100%. This finding is also in close agreement with that of Hebard et al.4 The curves also indicate a remarkably high Meissner fraction of about 50%, indicating that flux trapping is not very severe. Our observations in high magnetic fields are that the upper critical magnetic field slope, $-dH_{c2}/dT$, is quite steep and that the temperature of irreversibility is ~14 K in a magnetic

Conclusions. This work demonstrates that superconducting K_xC₆₀ can be synthesized by a much simpler solution chemistry route that avoids the use of vapor-phase K transfer and reaction. It is curious, however, that the bulk product seems to be similar in the extent of volume superconductivity ($\sim 1\%$) to that reported by Hebard et al.4 This suggests that the actual compound composition will become known only when highly crystalline singlephase material is synthesized and characterized. As such material is obtained, the Tc's will also likely increase significantly, especially as C₆₀ spheroid static^{13a} disorder below 77 K is suppressed.

Note Added in Proof. Contrary to the suggestion4 that the C60 spheroids should be in contact for an enhanced T_{c} , our experience with superconducting organic charge-transfer salts leads us to expect that T_c may actually increase with increase in the size of the alkali metal as it pries apart the C₅₀ spheroids and softens their interactions. 13b Thus, we have recently found that Rb $_{\rm c}$ C $_{60}$ prepared by the same solution-phase synthesis has $T_{\rm c}=28.6~{\rm K}^{.13c}$

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To whom correspondence should be addressed. Chemistry Division, Argonne National Laboratory

Materials Science Division, Argonne National Laboratory.

(17) University of California.

Chemistry and Materials Science Divisions, Argonne National Laboratory, Argonne, Illinois 60439, and Department of Chemistry, University of California, Irvine, California 92717

Aravinda M. Kini*,14-16 Brad M. Savall^{15,16} K. Douglas Carlson 15,16 Jack M. Williams*,14-16 Keith R. Lykke*,14-16 Peter Wurz^{15,16} Deborah Holmes Parker¹⁵⁻¹⁷ Michael J. Pellin*,14-16 Dieter M. Gruen^{15,16} Ulrich Welp*,14,16 Wai-Kwong Kwok*,14,16 Steven Fleshler¹⁶ George W. Crabtree¹⁶

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