C₆₀: Buckminsterfullerene

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During experiments aimed at understanding the mechanisms by which long-chain carbon molecules are formed in interstellar space and circumstellar shells¹, graphite has been vaporized by laser irradiation, producing a remarkably stable cluster consisting of 60 carbon atoms. Concerning the question of what kind of 60-carbon atom structure might give rise to a superstable species, we suggest a truncated icosahedron, a polygon with 60 vertices and 32 faces, 12 of which are pentagonal and 20 hexagonal. This object is commonly encountered as the football shown in Fig. 1. The C_{60} molecule which results when a carbon atom is placed at each vertex of this structure has all valences satisfied by two single bonds and one double bond, has many resonance structures, and appears to be aromatic.

The technique used to produce and detect this unusual molecule involves the vaporization of carbon species from the surface of a solid disk of graphite into a high-density helium flow, using a focused pulsed laser. The vaporization laser was the second harmonic of Q-switched Nd:YAG producing pulse energies of ~30 mJ. The resulting carbon clusters were expanded in a supersonic molecular beam, photoionized using an excimer laser, and detected by time-of-flight mass spectrometry. The vaporization chamber is shown in Fig. 2. In the experiment the pulsed valve was opened first and then the vaporization laser was fired after a precisely controlled delay. Carbon species were vaporized into the helium stream, cooled and partially equilibrated in the expansion, and travelled in the resulting molecular beam to the ionization region. The clusters were ionized by direct one-photon excitation with a carefully synchronized excimer laser pulse. The apparatus has been fully described previously²⁻⁵.

The vaporization of carbon has been studied previously in a very similar apparatus⁶. In that work clusters of up to 190 carbon atoms were observed and it was noted that for clusters of more than 40 atoms, only those containing an even number of atoms were observed. In the mass spectra displayed in ref. 6, the C_{60} peak is the largest for cluster sizes of >40 atoms, but it is not completely dominant. We have recently re-examined this system and found that under certain clustering conditions the C_{60} peak can be made about 40 times larger than neighbouring clusters.

Figure 3 shows a series of cluster distributions resulting from variations in the vaporization conditions evolving from a cluster distribution similar to that observed in ref. 3, to one in which C_{60} is totally dominant. In Fig. 3c, where the firing of the vaporization laser was delayed until most of the He pulse had passed, a roughly gaussian distribution of large, even-numbered clusters with 38-120 atoms resulted. The C₆₀ peak was largest but not dominant. In Fig. 3b, the vaporization laser was fired at the time of maximum helium density; the C₆₀ peak grew into a feature perhaps five times stronger than its neighbours, with the exception of C_{70} . In Fig. 3a, the conditions were similar to those in Fig. 3b but in addition the integrating cup depicted in Fig. 2 was added to increase the time between vaporization and expansion. The resulting cluster distribution is completely dominated by C₆₀, in fact more than 50% of the total large cluster abundance is accounted for by C_{60} ; the C_{70} peak has diminished in relative intensity compared with C₆₀, but remains rather prominent, accounting for ~5% of the large cluster population.

Our rationalization of these results is that in the laser vaporization, fragments are torn from the surface as pieces of the planar

Fig. 1 A football (in the United States, a soccerball) on Texas grass. The C₆₀ molecule featured in this letter is suggested to have the truncated icosahedral structure formed by replacing each vertex on the seams of such a ball by a carbon atom.



graphite fused six-membered ring structure. We believe that the distribution in Fig. 3c is fairly representative of the nascent distribution of larger ring fragments. When these hot ring clusters are left in contact with high-density helium, the clusters equilibrate by two- and three-body collisions towards the most stable species, which appears to be a unique cluster containing 60 atoms.

When one thinks in terms of the many fused-ring isomers with unsatisfied valences at the edges that would naturally arise from a graphite fragmentation, this result seems impossible: there is not much to choose between such isomers in terms of stability. If one tries to shift to a tetrahedral diamond structure, the entire surface of the cluster will be covered with unsatisfied valences. Thus a search was made for some other plausible structure which would satisfy all sp² valences. Only a spheroidal structure appears likely to satisfy this criterion, and thus Buckminster Fuller's studies were consulted (see, for example, ref. 7). An unusually beautiful (and probably unique) choice is the truncated icosahedron depicted in Fig. 1. As mentioned above, all valences are satisfied with this structure, and the molecule appears to be aromatic. The structure has the symmetry of the icosahedral group. The inner and outer surfaces are covered with a sea of π electrons. The diameter of this C_{60} molecule is ~7 Å, providing an inner cavity which appears to be capable of holding a variety of atoms8.

Assuming that our somewhat speculative structure is correct, there are a number of important ramifications arising from the existence of such a species. Because of its stability when formed under the most violent conditions, it may be widely distributed in the Universe. For example, it may be a major constituent of circumstellar shells with high carbon content. It is a feasible constituent of interstellar dust and a possible major site for

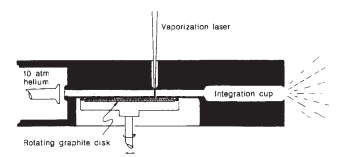


Fig. 2 Schematic diagram of the pulsed supersonic nozzle used to generate carbon cluster beams. The integrating cup can be removed at the indicated line. The vaporization laser beam (30-40 mJ at 532 nm in a 5-ns pulse) is focused through the nozzle, striking a graphite disk which is rotated slowly to produce a smooth vaporization surface. The pulsed nozzle passes high-density helium over this vaporization zone. This helium carrier gas provides the thermalizing collisions necessary to cool, react and cluster the species in the vaporized graphite plasma, and the wind necessary to carry the cluster products through the remainder of the nozzle. Free expansion of this cluster-laden gas at the end of the nozzle forms a supersonic beam which is probed 1.3 m downstream with a time-of-flight mass spectrometer.

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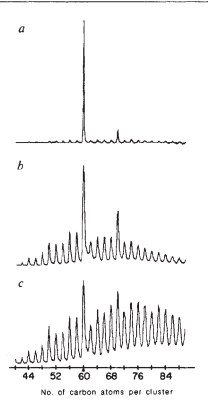


Fig. 3 Time-of-flight mass spectra of carbon clusters prepared by laser vaporization of graphite and cooled in a supersonic beam. Ionization was effected by direct one-photon excitation with an ArF excimer laser (6.4 eV, 1 mJ cm⁻²). The three spectra shown differ in the extent of helium collisions occurring in the supersonic nozzle. In c, the effective helium density over the graphite target was less than 10 torr-the observed cluster distribution here is believed to be due simply to pieces of the graphite sheet ejected in the primary vaporization process. The spectrum in b was obtained when roughly 760 torr helium was present over the graphite target at the time of laser vaporization. The enhancement of C₆₀ and C₇₀ is believed to be due to gas-phase reactions at these higher clustering conditions. The spectrum in a was obtained by maximizing these cluster thermalization and cluster-cluster reactions in the 'integration cup' shown in Fig. 2. The concentration of cluster species in the especially stable C₆₀ form is the prime experimental observation of this study.

surface-catalysed chemical processes which lead to the formation of interstellar molecules. Even more speculatively, C_{60} or a derivative might be the carrier of the diffuse interstellar lines⁹.

If a large-scale synthetic route to this C₆₀ species can be found, the chemical and practical value of the substance may prove extremely high. One can readily conceive of C₆₀ derivatives of many kinds—such as C₆₀ transition metal compounds, for example, C₆₀Fe or halogenated species like C₆₀F₆₀ which might be a super-lubricant. We also have evidence that an atom (such as lanthanum⁸ and oxygen¹) can be placed in the interior, producing molecules which may exhibit unusual properties. For example, the chemical shift in the NMR of the central atom should be remarkable because of the ring currents. If stable in macroscopic, condensed phases, this C₆₀ species would provide a topologically novel aromatic nucleus for new branches of organic and inorganic chemistry. Finally, this especially stable and symmetrical carbon structure provides a possible catalyst and/or intermediate to be considered in modelling prebiotic chemistry.

We are disturbed at the number of letters and syllables in the rather fanciful but highly appropriate name we have chosen in the title to refer to this C₆₀ species. For such a unique and centrally important molecular structure, a more concise name would be useful. A number of alternatives come to mind (for

example, ballene, spherene, soccerene, carbosoccer), but we prefer to let this issue of nomenclature be settled by consensus.

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- 1. Heath, J. R. et al. Astrophys. J. (submitted).
- 2. Dietz, T. G., Duncan, M. A., Powers, D. E. & Smalley, R. E. J. chem. Phys. 74, 6511-6512
- 3. Powers, D. E. et al. J. phys. Chem. 86, 2556-2560 (1982).
- Hopkins, J. B., Langridge-Smith, P. R. R., Morse, M. D. & Smalley, R. E. J. chem. Phys. 78, 1627-1637 (1983).
- 5. O'Brien, S. C. et al. J. chem. Phys. (submitted).
- 6. Rohlfing, E. A., Cox, D. M. & Kaldor, A. J. chem. Phys. 81, 3322-3330 (1984).
- 7. Marks, R. W. The Dymaxion World of Buckminster Fuller (Reinhold, New York, 1960).
- 8. Heath, J. R. et al. J. Am. chem. Soc. (in the press).
- 9. Herbig, E. Astrophys. J. 196, 129-160 (1975).

High-resolution solid-state NMR of quadrupolar nuclei

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Quadrupolar nuclei are the most abundant nuclear magnetic resonance (NMR)-receptive nuclei in the Earth's crust, and in many amorphous materials of technological interest (such as zeolite catalysts, ceramics and alloys), and have thus been intensively studied¹⁻⁷. Of particular interest is the ability to resolve and quantitate the various types of sites present in a given material. Here we present a very simple, yet we believe powerful, approach towards the resolution of chemically non-equivalent sites in solids, which combines a conventional high-field spin-echo NMR method with the resolution enhancement of the 'quadrupole shift' approach^{4,7}. We demonstrate its application to the complete resolution of both the (1/2, 3/2) and (3/2, 5/2) transitions of the ²⁷Al nuclei in a mixture of potassium and ammonium alums (KAl(SO₄)₂·12H₂O) and NH₄Al(SO₄)₂·12H₂O).

The major problem in spin-echo studies of most quadrupolar nuclei in solids is that a series of echoes is usually obtained^{8,9}. Solomon first reported⁸ the refocusing of first-order quadrupolar interactions using two radio-frequency pulses of the same phase, and Butterworth⁹ was able to differentiate quadrupolar from purely magnetic interactions in various alloys. However, this approach is not particularly useful for minerals and other nonmetallic materials, where large magnetic inhomogeneities are absent.

Fortunately, however, Bonera and Galimberti¹⁰ and Weisman and Bennett¹¹ showed that considerable enhancement of the 'desired' 2τ echo could be achieved by introducing a 90° phase-shift between the two radio-frequency pulses. Moreover, Weisman and Bennett showed (for nuclear spin I=5/2) for the special case of a $90_{0^{\circ}}-\tau-45_{90^{\circ}}$ pulse sequence that the amplitude of the 2τ echo (central plus quadrupolar contribution) was maximized, while the 'allowed' $3/2\tau$ plus 3τ echo amplitude was minimized¹¹. In practice, the 2τ echo is often the only one observed¹¹, and we show here that the Fourier transform of the echo can yield an essentially undistorted (1/2, -1/2; 1/2, 3/2; 3/2, 5/2) powder spectrum.

Figure 1a and b show the 23 Na and 27 Al spin-echo spectra of NaNO₃ and KAl(SO₄)₂·12H₂O, obtained at magnetic field strengths of 8.45 and 3.52 T, respectively, the response of the potassium alum being obtained in the presence of 1 H dipolar