Synthesis of C-59(CHPh(2))N from (C59N)(2) and C59HN. The first derivatization of C59N
Bellavia-Lund, C.; Gonzalez, R.; Hummelen, J.C.; Hicks, R.G.; Sastre, A.; Wudl, F.

Published in:
Journal of the American Chemical Society

DOI:
10.1021/ja964447c

IMPORTANT NOTE: You are advised to consult the publisher's version (publisher's PDF) if you wish to cite from it. Please check the document version below.

Document Version
Publisher's PDF, also known as Version of record

Publication date:
1997

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA):
Synthesis of C_{59}(CHPh_{2})N from (C_{30}N)_{2} and C_{39}HN. The First Derivatization of C_{39}N

Cheryl Bellavia-Lund, Rosario González, Jan Cornelis Hummelen, Robin G. Hicks, Angela Sastre, and Fred Wudl*  

Institute for Polymers and Organic Solids  
Departments of Chemistry and Materials  
University of California  
Santa Barbara, California 93106-5090  

Received December 30, 1996

Since the incorporation of heteroatoms into the fullerene skeleton is expected to modify its structural and electronic properties, we prepared the first heterofullerene, “C_{30}N”, which was isolated as its dimer, (C_{30}N)_{2}. The isolation in the condensed phase of this nitrogen-substituted fullerene allows for the investigation of the chemical reactivity, physical properties, and possible applications of this new class of fullerenes. As part of our continued interest in the azafullerenes, we have initiated studies to prepare derivatives. Even though N-labeling experiments and the isolation and full characterization of C_{39}HN have given us confidence that the starred carbons (see eq 1, below) are sp^{3} hybridized, a principal reason for derivatization of C_{39}N was to glean information on the variability of the chemical shift of the carbon atoms in question as a function of substitution.

Two possible processes for derivatization of C_{39}N are (1) deprotonation of the azafullerene C_{39}HN followed by attack of an electrophile and (2) homolytic dissociation of the interdimer bond of (C_{30}N)_{2} followed by free radical reactions. Here, we report on the derivatization of azafullerene via homolysis of its dimer.

According to theoretical calculations the interdimer bond of (C_{30}N)_{2} is relatively weak (18 kcal/mol) and should, under photolysis or thermolysis conditions, undergo facile homolysis. In a presence of a good hydrogen atom donor or radical source, the resulting azafullerenyl radical is expected to undergo free radical reactions.

Treatment of (C_{30}N)_{2} with excess diphenylmethane in refluxing o-dichlorobenzene (ODCB) for 48 h afforded, upon chromatography (silica gel, toluene), a crude product. HPLC analysis indicated one major constituent, consisting of 71 % of the isolable mixture. This was shown to be the substituted azafullerene C_{59}(CHPh_{2})N (2) by {sup 1}H and {sup 13}C NMR, FTIR, UV-vis, mass spectrometry, and cyclic voltammetry. The heterofullerene 2 was obtained in 42% isolated yield after purification by HPLC on a Cosmosil Buckypep semiprep column.

The {sup 1}H NMR spectrum of 2 showed signals corresponding to two equivalent phenyl groups and a singlet corresponding to the methine hydrogen at 6.09 ppm. The 34 lines observed between 156 and 124 ppm (30 from the C_{39}N moiety and 4 from the phenyl groups) in the {sup 13}C NMR spectrum are consistent with the expected C_{39}N symmetry of 2 depicted in eq 1. Another peak at 65.2 ppm corresponds to the methine carbon atom and a signal at 86.3 ppm must be due to the starred sp^{3} carbon atom on the buckyball moiety. This carbon resonance is downfield shifted by 14 ppm relative to the sp^{3} carbon of C_{39}HN, as expected on going from hydrogen to alkyl substitution. This is in agreement with a “closed” structure in which the sp^{3} carbon atom is bound to the nitrogen atom and the CHPh_{2} substituent, as shown in 2 above.

While the NOE effect identifies most of the phenyl resonances in the {sup 13}C NMR spectrum, proton-coupled {sup 13}C NMR distinguishes the resonance of the quaternary carbon of the phenyl group (quartet at 139 ppm) from the surrounding fullerene resonances. The coupled spectrum also supports the assignment of the methine carbon, revealing a doublet at 65.2 ppm (J_{CH} = 127 Hz) and a β-coupled doublet at 86.7 ppm (J_{CH} = 5 Hz) for the sp^{3} carbon of the fullerene cage. Finally, positive ion electrospray MS shows a molecular ion at m/z = 889 and a strong peak at m/z = 722 corresponding to C_{39}N.

The UV-vis absorption of C_{59}(PhCH)N is almost identical to that of C_{30}N and C_{39}HN showing a broad band in the visible at 448 nm; all three compounds are green in solution. The similarities of the electronic spectra of the azafullerenes supports the closed structure assigned to (C_{30}N)_{2} depicted for 1, above.

The FTIR spectrum shows a typical pattern for a fullerene derivative, in particular having absorptions similar to C_{39}HN in the fingerprint region (480-590 cm^{-1}), where the strongest peak is at 529.3 cm^{-1}.

The diphenylmethane adduct of C_{39}N exhibits two quasi-reversible one-electron reduction waves; a third wave, which is completely irreversible was also observed. This contrasts the behavior of C_{39}HN in which the reduction processes are chemically irreversible. We interpreted the properties of the latter as arising from facile hydrogen atom loss on reduction, which is not possible in the diphenylmethane derivative. The reduction potentials are presented in Table 1, along with those of C_{30}N, (C_{30}N)_{2}, and C_{39}HN for comparison. The potentials of the (diphenylmethyl)azafullerene and hydroxoaazafullerene are comparable, and both are significantly more difficult to reduce than the dimer, an observation consistent with the high electronegativity of a heterofullerene “substituent”. All of the azafullerenic compounds possess irreversible oxidation waves at much lower potentials compared to those of C_{60}.


(8) (PhCH)_{2}N: Diphenylmethane (2.4 mL) was added to a degassed solution of (C_{30}N)_{2} (30 mg) in HPLC grade 1,2-dichlorobenzene (25 mL). The mixture was refluxed under argon for 48 h and chromatographed on silica gel using toluene as eluent. The product was purified by HPLC using a semipreparative Cosmosil Buckypep column (30% hexane in toluene, 5 mL/min, UV/326 nm). Solvents were removed under vacuum, and the product was washed with ether, centrifuged, and decanted three times to remove ether-soluble components, including toluene. It was finally dried under vacuum. Yield: 15 mg (42%). Please refer to Supporting Information for spectroscopic data.

S0002-7863(96)04447-2 CCC: $14.00 © 1997 American Chemical Society
### Table 1. Redox Potentials for C\textsubscript{60} and some C\textsubscript{59}N Derivatives\textsuperscript{a}

<table>
<thead>
<tr>
<th></th>
<th>C\textsubscript{60}</th>
<th>(C\textsubscript{59}N)\textsubscript{2}</th>
<th>C\textsubscript{59}HN</th>
<th>C\textsubscript{59}N(CH\textsubscript{2}Ph\textsubscript{2})</th>
</tr>
</thead>
<tbody>
<tr>
<td>$E_{1}^{\text{red}}$</td>
<td>-1123</td>
<td>-992</td>
<td>-1106\textsuperscript{b}</td>
<td>-1082</td>
</tr>
<tr>
<td>$E_{2}^{\text{red}}$</td>
<td>-1450</td>
<td>-1471</td>
<td>-1485</td>
<td></td>
</tr>
<tr>
<td>$E_{3}^{\text{red}}$</td>
<td>-1913</td>
<td>-1979</td>
<td>-2089</td>
<td></td>
</tr>
<tr>
<td>$E_{\text{ox}}^{c}$</td>
<td>+1300\textsuperscript{b,c}</td>
<td>+886</td>
<td>+823\textsuperscript{b}</td>
<td>+867\textsuperscript{b}</td>
</tr>
</tbody>
</table>

\textsuperscript{a} All potentials are in mV vs Ferrocene, in ODCB unless stated otherwise. \textsuperscript{b} Chemically irreversible. \textsuperscript{c} Dubois, D.; Kadish, K. J. Am. Chem. Soc. 1991, 113, 7773. \textsuperscript{d} Chemically reversible, in 1,1,2,2-tetrachloroethane solvent: Xie, Q.; Arias, F.; Echegoyen, L. J. Am. Chem. Soc. 1993, 115, 9818.

We can attribute the mechanism of formation of 2 to be a short free radical chain. As illustrated in eq 2, the initiation step is proposed to be the homolysis of the azafullerene dimer to C\textsubscript{59}N\textsuperscript{•}.

\[
(C_{59}N)_2 \xrightarrow{\text{ODCB}} 2C_{59}N^\cdot \quad (2)
\]

Subsequently, the steps in eqs 3a and 3b are plausible propagation and termination steps. Hence, in the presence of the hydrogen donor diphenylmethane, formation of C\textsubscript{59}HN was expected to be one of the major products (eq 3a); however, it was only 2% of the reaction mixture. The major product was the diphenylmethane derivative of the azafullerene radical, 2, (eq 3b).

\[
C_{59}N^\cdot + \text{Ph}_2\text{CH}_2 \xrightarrow{\text{ODCB}} C_{59}HN + \text{Ph}_2\text{CH}^\cdot \quad (3a)
\]

\[
\text{Ph}_2\text{CH}^\cdot + C_{59}N^\cdot \xrightarrow{\text{ODCB}} 2 \quad (3b)
\]

Two possible reasons for the low yield of C\textsubscript{59}HN are (1) unlike 2, which appears to be quite stable under the “harsh” reaction conditions, C\textsubscript{59}HN decomposes or polymerizes to yield insoluble products or (2) C\textsubscript{59}HN undergoes a free radical reaction to afford C\textsubscript{59}N\textsuperscript{•} (eq 4). In eq 4, “In” under the arrow could be an adventitious initiator or C\textsubscript{59}N\textsuperscript{•} from (C\textsubscript{59}N)\textsubscript{2} (see eq 5, below).

\[
\begin{align*}
C_{59}NH & \xrightarrow{\text{In}^\cdot} C_{59}N^\cdot + \text{In}^- \\
2C_{59}N^\cdot & \xrightarrow{\text{ODCB}} (C_{59}N)_2 + \text{H}_2 \quad (4)
\end{align*}
\]

In a control experiment, treatment of C\textsubscript{59}HN\textsuperscript{•} with diphenylmethane in refluxing ODCB for 24 h, afforded, upon chromatography (silica gel, toluene), one isolable band, HPLC analysis of the crude product revealed the presence of 2 (73%), (C\textsubscript{59}N)\textsubscript{2} (7%), and unreacted C\textsubscript{59}N (5%). This experiment proves that equation 4 is operative and that In\textsuperscript{•} is due to a combination of eqs 5 and 2.

\[
\begin{align*}
2C_{59}N^\cdot & \xrightarrow{\text{ODCB}} (C_{59}N)_2 + \text{H}_2 \quad (5)
\end{align*}
\]

In support of the above dimerization, fresh, HPLC single component samples of C\textsubscript{59}HN were found to dimerize on standing at room temperature even in the solid state.

We have demonstrated above the relatively clean conversion of (C\textsubscript{59}N)\textsubscript{2} to a stable, soluble derivative in good yield. This study shows that C\textsubscript{59}HN can also undergo homolytic cleavage to yield the C\textsubscript{59}N radical, where all three species C\textsubscript{59}N\textsuperscript{•}, C\textsubscript{59}HN, and (C\textsubscript{59}N)\textsubscript{2} are involved in the process leading to 2. Full characterization of the title molecule shows remarkable similarities to (C\textsubscript{59}N)\textsubscript{2} and C\textsubscript{59}HN, and in particular the UV-vis data demonstrate that the electronic structure of all three molecules is essentially identical. The $^{13}$C NMR spectra reveal a similar pattern in the $sp^2$ region of 2 to that of its precursors and an unmistakable $sp^3$ carbon at 86.1 ppm, illustrating the chemical shift deviation as a result of substitution; an important stepping stone for the absolute determination of the (C\textsubscript{59}N)\textsubscript{2} structure.\textsuperscript{9} Research is currently in progress utilizing this methodology to derive azafullerenes with unusual properties.

### Acknowledgment

We thank the National Science Foundation for support through the Materials Research Laboratory and NSF DMR 95-00888, as well as individual investigator grants. R.G. and A.S. thank the Ministerio de Educación y Cultura (Spain) for postdoctoral fellowships. R.G.H. thanks the National Research Council of Canada for a postdoctoral fellowship.

### Supporting Information Available

Experimental details for the preparation, purification, and characterization of 2 (6 pages). See any current masthead page for ordering and Internet access instructions.

JA964447C