Research Article

Effect of Structure of Monocyclic Aromatic Solvents on the Packing Density of Fullerene C₆₀ Solvation Shell

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Fullerene C₆₀ solutions in various monocyclic aromatic solvents (AS) diluted by tetrahydrofurane (THF) were studied using UV spectroscopy. An algorithm was proposed which made it possible to obtain the data from the UV spectra of (C₆₀+AS) solutions in THF which pointed to the difference between the packing density of AS molecules in solvation shell of C₆₀ and that in the pure AS solvent. The presence of asymmetrical substituents in the AS molecule has been shown to result in loosening of solvent structure in the solvation shell, while the absence of the substituents or presence of symmetrical substituents leads to compacting of solvent structure.

1. Introduction

Donor-acceptor interactions of fullerene C₆₀ with solvent molecules are responsible for its good solubility in numerous organic solvents due to formation of strong solvation shells [1]. Fullerene C₆₀ manifests itself mainly as a π-acceptor [1–3], which predominates its high solubility in aromatic solvents due to formation of fullerene-solvent molecular complexes. The presence of substituents in the aromatic solvent (AS) molecule may be expected to result in the appearance of additional interactions between the AS and C₆₀ during solvation. For example, the spectroscopic study of fullerene solutions in various solvents [4, 5] pointed to the possibility of C₆₀ solvation by o-dichlorobenzene owing to both π–π interactions of the benzene ring of AS with the π-system of fullerene and partial charge transfer from the chlorine atom to fullerene. The presence of methyl group as a substituent in the AS molecule may open the way to formation of additional hydrogen-bonding complexes [6]. Interaction of some organic donors with C₆₀ has been shown to result in their coordination with fullerene C₆₀ [7]. The structure of solvation shell would be dependent on the forced coordination of solvent molecules by fullerene. The fullerene ability to structure a solvent was shown in [8]. It seems obvious that the coordination of monocyclic AS molecules by C₆₀ during solvation would depend on the presence of substituents in AS which may interact with fullerene. The forced coordination of solvent molecules by C₆₀ may be proposed to result in the difference between the packing density of AS molecules in the C₆₀ solvation shell and that in the pure AS. The examination of this idea was the aim of the work.

2. Experimental Part

2.1. Materials, Experimental Procedure, and Equipment. Fullerene C₆₀ (purity 99.5 wt%) (Fullerene Technologies, St. Petersburg) was used for preparation of its solutions with specified concentrations in several selected aromatic solvents with various substituents (Table 1) as well as in tetrahydrofurane (THF) and methylene chloride (CH₂Cl₂). All solvents were preliminary purified by distillation.

The weighed amount of C₆₀ solution in AS with the known fullerene concentration was diluted by the weighed amount of THF, which allowed us to calculate the concentration of fullerene (c_F) and of the AS used (c_AS,cal) in the (C₆₀+AS) solution in THF. No less than three solutions diluted by THF were prepared from the C₆₀ solution in each AS;
Table 1: Aromatic solvents used and the number $N$ of excess (positive values) or deficient (negative values) molecules of the aromatic solvent per solvation shell of C$_{60}$ molecule.

<table>
<thead>
<tr>
<th>Solvent</th>
<th>Structural formula</th>
<th>$N$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benzene</td>
<td><img src="image" alt="Benzene" /></td>
<td>+650</td>
</tr>
<tr>
<td>Toluene</td>
<td><img src="image" alt="Toluene" /></td>
<td>−110</td>
</tr>
<tr>
<td>o-xylene</td>
<td><img src="image" alt="o-xylene" /></td>
<td>−240</td>
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<tr>
<td>p-xylene</td>
<td><img src="image" alt="p-xylene" /></td>
<td>+940</td>
</tr>
<tr>
<td>Chlorobenzene</td>
<td><img src="image" alt="Chlorobenzene" /></td>
<td>−190</td>
</tr>
<tr>
<td>o-dichlorobenzene</td>
<td><img src="image" alt="o-dichlorobenzene" /></td>
<td>−120</td>
</tr>
</tbody>
</table>

for each (C$_{60}$+AS) solution in THF the UV spectrum was recorded.

The UV spectra of C$_{60}$ solutions were recorded using a UV VIS SF-2000 spectrophotometer (Russia); quartz cells of 2 mm thickness were used.

2.2. The Procedure of Calculation of the Number of Excess or Deficient Molecules of AS per Solvation Shell of C$_{60}$ Molecule. Tetrahydrofuran is a transparent in the UV-region solvent good for AS and poor for C$_{60}$. At addition of THF to C$_{60}$ solution in AS, fullerene solvation shell will be defined as the number of AS molecules which C$_{60}$ is able to retain in THF. If the packing density of AS molecules in solvation shell corresponds to that in the pure AS solvent, the concentration $c_{AS,sp}$ calculated from the spectra of the (C$_{60}$+AS) solution in THF must be equal to the concentration $c_{AS,cal}$ calculated for this solution from the weighed amounts of C$_{60}$ solution in AS and of THF. To calculate $c_{AS,sp}$ concentration, it is necessary to isolate the C$_{60}$ spectrum from the (C$_{60}$+AS) spectrum in the region of the combined absorbance of AS and C$_{60}$ (at $\lambda < 300$ nm). The following algorithm was developed to evaluate the $c_{AS,sp}$ concentrations of AS from the spectra of (C$_{60}$+AS) solutions in THF.

The graphic illustration of the algorithm proposed is presented in Figure 1. Curve 1 in Figure 1 corresponds to the spectrum of (C$_{60}$+AS) solution in THF with known concentration $c_{AS,cal}$ and, supposedly, is a superposition of C$_{60}$ and AS spectra.

To recover the individual C$_{60}$ spectrum, the following calibration procedure was developed. From the UV spectra recorded for C$_{60}$ solution in CH$_2$Cl$_2$ under gradual addition of THF (Figure 2), the linear dependence, $I_A F = C \cdot A_F$, of the integral absorbance $I_A F$ of C$_{60}$ (the area under every
The insert shows the normalized UV spectra of (1) C60 in THF, (2) (C60+CH2Cl2) in THF, and (C60+AS) solutions in THF in the region transparent for C60: (3) AS is toluene, (4) AS is p-xylene, and (5) AS is chlorobenzene.

The UV spectra of C60 (1) in THF and (2) in CH2Cl2. Curve 2 in Figure 1 corresponds to the C60 spectrum obtained by transformation of the spectrum of (C60 + CH2Cl2) solution in THF with the known value of integral absorbance (for any spectral curve in Figure 2). The transformation of the selected standard C60 spectrum with the known IA_F value to the C60 spectrum corresponding to the absorbance value at λ_max = 329.3 nm in the (C60+AS) spectrum was carried out by multiplication of absorbance values of the standard C60 spectrum by the ratio of integral absorbance calculated from the calibration dependence IA_F = C ∙ A_F to the integral absorbance of the standard spectrum.

From the UV spectra of AS solutions in THF recorded at specified concentrations c_AS of AS (Figure 4) the linear calibration dependence IA_AS = B_AS ∙ c_AS (the insert in Figure 4) were obtained for all aromatic solvents used, where IA_AS is the integral absorbance (the area under the spectral curve of AS at c_AS) within the absorption range of AS (from λ_1 to λ_2, Figure 4).

Using the calibration IA_AS = B_AS ∙ c_AS, the standard spectrum of AS (the spectrum with a known value of IA_AS) in THF was transformed to the AS spectrum corresponding to the respective concentration c_AS,cal of AS in the (C60+AS) solution in THF (curve 3 in Figure 1). The spectrum (curve 4 in Figure 1) obtained by subtraction of the absorbance values of C60 (curve 2) from the absorbance values of (C60+AS) (curve 1) is the spectrum of AS which corresponds to the true concentration c_AS,sp of AS in solution. The integral absorbance IA_AS,sp of AS was obtained as the difference between the integral absorbance of (C60+AS) solution and the calculated integral absorbance IA_F of C60. The concentration c_AS,sp of AS was calculated from the value of IA_AS,sp, obtained from the calibration IA_AS = B_AS ∙ c_AS.

In accordance with the c_AS,cal and c_AS,sp values obtained for each (C60+AS) solution in THF; the number N of excess (positive values) or deficient (negative values) molecules of AS per solvation shell of C60 molecule was calculated from the relation

\[ N = \frac{\Delta c_{AS}}{c_F - c_{AS,cal}} \]

where \( \Delta c_{AS} = (c_{AS,sp} - c_{AS,cal}) \), \( c_F \), and \( c_{AS} \) are the molecular masses of C60 and AS, respectively.

3. Results and Discussion

For all (C60+AS) solutions in THF the AS concentration c_AS,sp obtained with the proposed algorithm differed from the c_AS,cal value. Thus, the comparison of the chlorobenzene spectrum corresponding to the calculated concentration c_chlorobenzene,cal in THF (curve 3 in Figure 5) with the spectrum corresponding to the true concentration of chlorobenzene (curve 4 in Figure 5) shows that the true concentration is lower than the calculated one.

The comparison of the spectrum corresponding to the calculated p-xylene concentration c_p-xylene,cal (curve 3 in Figure 6) with the spectrum corresponding to the true concentration of p-xylene (curve 4 in Figure 6) shows that...
the true concentration exceeds considerably the calculated concentration.

The difference between the $c_{AS,cal}$ and $c_{AS,sp}$ values found for each (C$_{60}$+AS) solution in THF points to the change of solvation shell density with respect to the density of pure solvent. The calculated $N$ values are given in Table 1. The data show that the presence of asymmetrical substituents (chlorine atoms or the methyl group) in the AS molecule results in the loosening of solvent structure in the solvation shell of C$_{60}$, while the absence of substituents (benzene) or the presence of symmetrical substituents (p-xylene) leads to compacting of solvent structure in the C$_{60}$ solvation shell. The insignificant spread in the calculated $N$ values (below 10%) supports the validity of the proposed algorithm for calculation of the true AS concentration from the UV spectroscopic data. The validity of the proposed algorithm is supported as well by the practical identity of the normalized real spectra of AS in THF and of the AS spectra obtained as a difference of spectra of (C$_{60}$+AS) and C$_{60}$ (see inserts in Figures 5 and 6). The concentration dependences of the integral absorbance of C$_{60}$ in various aromatic solvents (Figure 7) correlate with the data listed in Table 1. Absorption decreases with the increase of packing density of solvent molecules in the solvation shell of C$_{60}$.

4. Conclusions

The idea has been examined that the forced coordination of monocyclic molecules with various substituents of aromatic solvents by fullerene C$_{60}$ may result in the difference between

Figure 5: The UV spectra of (1) (C$_{60}$+o-dichlorobenzene) solution in THF ($c_{o-dichlorobenzene,cal}$ = 2.113 mg/mL and $c_F$ = 0.0095 mg/mL), (2) the transformed standard spectrum of C$_{60}$, (3) o-dichlorobenzene corresponding to $c_{o-dichlorobenzene,cal}$ = 2.113 mg/mL (IA$_{o-dichlorobenzene}$ = 9.598 $\cdot$ C$_{60}$-dichlorobenzene in the $\lambda$ range from 245 to 285 nm), and (4) o-dichlorobenzene obtained as a difference of (1) and (2) spectra. The calculated value $c_{o-dichlorobenzene,sp}$ for the spectrum (4) is 1.851 mg/mL. The insert shows the normalized spectrum of C$_{60}$ in THF (solid line) and the normalized spectra of o-dichlorobenzene obtained by subtraction of the C$_{60}$ spectrum from the spectra of (C$_{60}$+o-dichlorobenzene) solutions in THF with various concentrations of o-dichlorobenzene and C$_{60}$: $c_{o-dichlorobenzene,cal}$ = 2.113 mg/mL, $c_F$ = 0.0095 mg/mL (dash line); $c_{o-dichlorobenzene,cal}$ = 2.190 mg/mL, $c_F$ = 0.0069 mg/mL (short dash line); $c_{o-dichlorobenzene,cal}$ = 1.758 mg/mL, $c_F$ = 0.0081 mg/mL (dot line).

Figure 6: The UV spectra of (1) (C$_{60}$+p-xylene) solution in THF ($c_{p-xylene,cal}$ = 1.548 mg/mL and $c_F$ = 0.0061 mg/mL), (2) the transformed standard spectrum of C$_{60}$, (3) p-xylene in THF corresponding to $c_{p-xylene}$ = 1.548 mg/mL (IA$_{p-xylene}$ = 9.013 $\cdot$ C$_{60}$-p-xylene in the $\lambda$ range from 240 to 280 nm), and (4) p-xylene obtained as a difference of (1) and (2) spectra (with the calculated value of $c_{p-xylene,sp}$ = 2.495 mg/mL). The insert shows the normalized spectrum of p-xylene in THF (solid line) and normalized spectra of p-xylene obtained by subtraction of the C$_{60}$ spectrum from the spectra of (C$_{60}$+p-xylene) solutions in THF with various concentrations of p-xylene and C$_{60}$: $c_{p-xylene,cal}$ = 1.548 mg/mL, $c_F$ = 0.0061 mg/mL (dash line); $c_{p-xylene,cal}$ = 1.166 mg/mL, $c_F$ = 0.0063 mg/mL (short dash line); $c_{p-xylene,cal}$ = 1.378 mg/mL, $c_F$ = 0.0070 mg/mL (dot line).

Figure 7: Concentration dependences of the integral absorbance (in the $\lambda$ range from 300 to 365.1 nm) of C$_{60}$ in (1) o-xylene, (2) toluene, and (3) p-xylene.
the packing density of AS molecules in the $C_{60}$ solvation shell and that in the pure AS. In this work, the fullerene solvation shell was defined as the number of AS molecules which the molecule of fullerene is able to retain at addition of THF which is a good solvent for AS and poor for $C_{60}$, to $C_{60}$ solution in AS. The analytical techniques applied for preparation of $C_{60}$ solutions in aromatic solvents and for determination of accurate concentrations $c_F$ and $c_{AS,cal}$ after addition of THF, as well as the developed calibration procedures, made it possible to obtain the data from the UV spectra of $(C_{60}+AS)$ solutions in THF which is the supporting evidence of the difference between the packing density of AS molecules in the solvation shell of $C_{60}$ and that in the pure AS solvent. The data obtained show that the presence of asymmetrical substituents (chlorine atoms or methyl group) in the AS molecule results in the loosening of solvent structure in the solvation shell of $C_{60}$, while the absence of substituents (benzene) or the presence of symmetrical substituents ($p$-xylene) leads to compacting of solvent structure in the $C_{60}$ solvation shell.

References


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