THE ARC DISCHARGE WITH A DIRECTIONAL GAS FLOW: SYNTHESIS AND PROPERTIES OF FULLERENES

M. M. Kasumov, V. Ya. Chernyak*

Taras Shevchenko National University of Kyiv, Prospect Glyshkova 4G, Kyiv, Ukraine

ABSTRACT. Discharge is ignited in a hollow electrode with a directional stream of gas. Under these conditions, the duration of fullerene formation in space at high density and temperature is longer, in product the amorphous component is not manifested and the heavy fullerene yield increases. There are the technique and the results of measuring the electro options. It noted the dependence of the spectrum of the charge of fullerenes and the preferred use of ions fullerenes.

KEYWORDS: amorphous component, electron affinity, hollow electrode, ionization potential, formation duration, gas-plasma stream, neutralizing of fullerenes, turbulence.

INTRODUCTION

General provisions

In the 1970s, the possibility of the existence of hollow C_{20}, C_{60} clusters and their electronic structures were shown theoretically [1]. These carbon clusters, which were called fullerenes, were obtained experimentally in a supersonic helium stream by the action of a laser beam on the graphite surface [2]. Investigations showed that fullerenes are effective electron acceptors and form compounds with new properties with atoms of other elements. The discovered peculiarities show fullerenes and their derivatives to be a new class of molecules-compounds; the trends in physics, chemistry, biology, medicine and technology, which are originating on their basis, are also new.

The most important among such is a composite organic semiconductors with fullerenes C_{60} which was used in a photoelectric converter [3]. The perfecting of a composite technology allowed to reach of solar cells efficiency to 6,7%[4], that is many times higher than in earlier known elements of the GaAs, ZnO, TiO_2 [5].

The big hopes are tied with water-soluble form of fullerenes, which allow you to enter the sphere of the use of different salt solutions including the lymph and blood person, and directly related to determining the health of the person. In ongoing studies [6], to develop methods to obtain more stable form of output. Therefore it was intended to carry out a technological revolution by modernizing the main spheres of activity through the use of fullerenes, their derivatives and composites based on them as the main material. To realize this idea, a search for and development of fullerene synthesis methods were initiated. At the present time, the most commonly used fullerene synthesis methods are: laser [2], chemical [7] and arc discharge [8] methods. The common peculiarity of the methods is that fullerene synthesis takes place in an inert gaseous medium at the temperature 1500 < T < 7000 K [9] (low - temperature plasma conditions). The formation of nanostructures in a space with high
temperature takes place by collision of carbon atoms and clusters (C₂, C₃, ...). The process is affected by the Coulomb interaction of particles with opposite sign. The synthesis methods differ in fullerene formation mechanism, but the results of the last few decades have shown that methods within the measurement error have a limit value output \( \eta = (19 \pm 4)\% \). The development of laser and chemical methods probably will eliminate the noted deficiencies [10]. But arc plasma medium on the basis of carbon vapor is an indispensable element to maintain the continuity of the arc and the synthesis process. Discharge control elements are: the geometry of the electrode, gas atmosphere and arc discharge parameters. An additional advantage of the method is the experience of creating and using a wide range of bit devices and the equipment for arc discharge.

The aim of this work is a more detailed discussion of the features of the new method of arc synthesis of fullerenes, which showed a significant advantage in comparison with the approved methods, and to study insufficiently known the electromagnetic characteristics and determine the electrical parameters of fullerenes.

The content of this article is in addition to [11], and it has been obtained mostly on the basis of a detailed analysis of the material in this work. So in article are kept the main peculiarities of the discharge device and formation process; to them are added the revealed in this material properties of positively charged fullerenes.

**On the Fullerene Formation Process in Arc Discharge**

Fullerene arc synthesis is a multievent process, the theory of which has not been carried to completion. According to general concepts, the \( C_j \) fullerene formation process takes place in the discharge space at 1500 – 5000 K during diffusion as a result of collisions followed by coalescence of nanoparticles. Fullerene synthesis is conventionally represented as series of acts of attachment of small \( C_k \), \( C_i \) clusters of carbon vapor from the heated portions of the electrodes and the subsequent kinetic excess energy relaxation of coalescing particles in inverse processes, annealing-away of \( n_f C_f \) clusters. Conventional synthesis scheme:

\[
C_j = C_k + \sum_i n_i C_i - \sum_f n_f C_f,
\]

where \( C_i \) is the initial fullerene cluster formed in the interelectrode space, the coefficients \( n_i \), \( n_f \) are integers. The clusters in the scheme (1) are summed with respect to the subscripts \( i \) and \( f \). The fullerene formation parameters are formed from the sum of individual segments: duration of the formation process, \( T_j = \sum_i t_{C_i} \), and formation path length \( H_j = \sum_i h_{C_i} \). During the formation stability of the structure is controlled by the annealing process, which is under the influence of atomic bonds at conventional scheme (1) correspond to particles \( \sum_f n_f C_f \). Structure formed with a number of carbon atoms believe resonance (C₆₀, C₇₀, ...), because the mechanism of formation of these structures is unknown. In arc discharge, the diffusion nature of spatial motion with temperature decrease from radius gives a spread of clusters in path parameters, in mass and quality of the product. In arc discharge, amorphous carbon black accounts for a larger fraction of the product (>50%). From the analysis of the existing concepts it may be assumed that the product of other quality can be obtained in a space with high temperature, in which longer duration and length of fullerene formation path with turbulent portions are ensured. The space with such peculiarities has a circular symmetry at the minimal energy consumption.
Problem Solution

Arc Discharge in a Hollow Electrode with Working Gas Flow (ADHE-WGF)

To ensure continuous synthesis process, a stock-produced TDM-317 as welding power source with dropping current-voltage characteristic and a pulsed arcing stabilizer (PAS) were used jointly. The TDM-317 + PAS circuit continuously holds plasma in the interelectrode space, also at the moments of passage of current and voltage through zero [12]. Presented further design of the discharge device was created based on the original dorobotki devices [13, 14]. In the works [15, 16], fullerene synthesis was carried out in a chamber of 68 mm diameter (h = 150 mm) on the discharge device shown in Fig. 1: graphite electrode 1 for emission spectrum analysis (diameter 6 mm), graphite electrode 2 (cavity diameter 15 mm, h = 15 mm) with slits through which working gas (He) enters the electrode cavity. The cuts (slots) in transverse and longitudinal dimensions provide education directed jet passing gas. In the design, the part 3 is a graphite cover. Electrical insulation between the electrodes 1 and 2 is provided by a high-temperature ceramic, an alund crucible 4. An extra function of crucible is heating the working gas that enters the discharge space. The working gas coming from the free space heats on the passage of gas-plasma flow and discharge radiation over the surface of the crucible, which heats up on thermal contact with the heated electrode portion. The parts 5 and 6 of common steel apply potential to the electrode 2 and fasten the construction together into a whole. The part 7 is the chamber wall.

On the Product of Synthesis by Arc Discharge in a ADHE-WGF

The composition of the product of synthesis by ADHE-WGF (fullerite) was determined on an AutoFlex apparatus (Bruker, Germany) from a time-of-flight mass spectrum (MS), Fig. 2. This is a MS of fullerenes with negative charge for a product obtained in ADHE-WGF under optimal conditions. As can be seen, the lines in this series follow at the interval ∆ (m/z) = 24 amu = 2m₄, which is usual for fullerene spectrum. It is seen from Fig. 2 that the MS consists of background bands, which differ in height and width, bands of medium hydrogenated clusters (C₂₆, C₂₇, C₂₈), C₆₀, C₇₀, C₇₂ and a series of C₇₄₂ₙ lines, where n = 0-40. In the spectrum, the lines of stable nanostructures, which accumulated during discharge on the cooled walls of the chamber: C₆₀, C₇₀,…, C₇₈, C₈₄,…, C₉₀, C₉₆, C₁₁₂,…, C₁₅₀ stand out in the value of the peaks. The peak values in the MS of fullerenes in Fig 2 are in agreement with the formation mechanism of scheme (1) and another detailed presentation proposed: the peak values of fullerenes decrease with increasing mass, but the decrease in peak values in ADHE-WGF is smaller as compared with ordinary-arc discharge spectrum. An important peculiarity of the spectrum of ADHE-WGF, which contradicts the above one, is that the peak value of C₈₄ fullerene exceeds that of the C₇₀ fullerene. The spectrum of fullerenes with positive charge ends with the C₈₄⁺ fullerene peak (see below), but the background spectrum has an extension at m/z > 1800 u.

On the Mechanism of Fullerene Formation in ADHE-WGF

As was pointed out above (1), fullerenes begin to form on collisions of carbon atoms, ions and small clusters C₂, …, C₄ by the action of a turbulent evaporation flow from the surface of the electrodes. But in a hollow electrode, the turbulence under the action of working gas flow is higher, and the process occurs with the participation of helium atoms via forward and back reactions [17]. Another distinction of the MS of ADHE-WGF from ordinary-discharge MS is background. The structures that create a background in the MS of ADHE-WGF don’t affect
the transparency of the benzene solution of fullerene carbon black. The background is created by nanostructures formed as a result of “re-formation” of fullerenes and annealing away of fragments, which become just a “nanobackground”. The nanobackground peaks in the mass spectrum $24 \leq m/z < 2000$ u have a value of up to $2.0 - 3.5 \%$ of the peak value of $C_{60}$. The MS of the product of ADHE-WGF at the discharge current $I = 70$ A without filtration coincides with the spectrum obtained after double filtration. The result is confirmed by the MS of fullerene carbon black synthesized at $I = 80$ A. There is a background in the spectrum of positive and negative ions, and the background is larger for the product obtained at higher discharge current.

**Geometrical Factor in Fullerene Formation Shows the Nature of the Limit Value Outputs**

In arc discharge, the length of fullerene formation path as a rule is estimated from the equation:

$$L_0 = (1 \pm 3) \times \Delta, \quad (2)$$

where \(\Delta\) is working interelectrode gap \(\Delta = 1 - 7 \text{ mm}\) [18], and the length of fullerene formation path \(L_0 \approx 15 \text{ mm}\).

An examination of the formation space in the device for ADHE-WGF is given on the basis of the size of the electrodes and gas plasma thermodynamics, gives the path:

$$L_1^* \geq 100 \text{ mm}. \quad (3)$$

In this case the trajectory of the fullerene formation shaped like a helix, which has 2-3 turns. Comparison of the values in (2) and (3) shows an advantage of the method with hollow electrode over that with ordinary discharge: *the fullerene yield is higher at the longer length of nonlaminar formation path.*

**The Measurement Results and Discussion**

The synthesized in ADHE-WGF product was analyzed in a solid fullerite phase on an AUTOFLEX RIIILRF 20 time-of-flight mass spectrometer with pulsed $N_2$ laser (Germany). The mass spectrum negatively charged fullerenes presented in Fig 2. The spectrum was obtained at the discharge current $I = 70$ A. The values of the peaks of the fullerenes $C_{60}$, $C_{70}$, $C_{84}$, ..., $C_{150}$ in the mass spectrum are as follows: 100, 57, 78, ..., 6. The descent of the envelope curve of the peaks of fullerene mass spectrum with increasing number of atoms in the cluster is common. This is in accordance with the generalized scheme (1). In discharge with hollow electrode, however, the decrease in the values of peaks is smaller, and the value peak $C_{84}$ is larger than peak $C_{70}$. The peculiarity apparently shows that the structure of the $C_{84}$ cluster is formed by the optimal number of covalent electron bonds, which have a three-dimensional structure. As a result, the $C_{84}$ cluster has been obtained, which is stronger than other fullerenes Fig 2. In mass spectrum is distinguished peaks lines (conditional resonant structures) poorly known fullerene $C_{112}$, $C_{122}$, $C_{146}$, $C_{150}$. To determine the values of the electron affinity (EA$_n$) and energy of ionization (EI$_n$) of fullerenes were made calculations. The calculations are based on a rigid model for the structure of the carbon spheres formed from $n$ carbon atoms with an effective radius of the $R_n = k\cdot n^{0.5}$. The calculations used in a ratio of adjusted:
\[ IE_n = \varphi_\infty - K_1/R_n \quad (4) \]

\[ and \quad EA_n = \varphi_\infty + K_2/R_n \quad , \quad (5) \]

\( n \) – integer of carbon atoms, \( K_1 \) and \( K_2 \) – normalization coefficients, \( \varphi_\infty = 5.37 \text{ eV} \) – the work of function for graphite sheet, \( k \) – coefficient for adjusting. Data for \( C_{60} \) were used in the calculations [18]. The results of calculations for noted of heavy fullerenes Fig. 2 are given in the Table.

According Table seen that the ionization energy of clusters decreases with increasing number carbon atoms, but with the number of carbon atoms increases the electron affinity. Therefore, according to Table between electro-physical parameters of fullerene \( C_n \) there is a general relationship [18]:

\[ EA_n + IE_n \leq 2\varphi_\infty \quad (6) \]

The values included in the relation (6) is by definition have a different nature, and show that the mass spectra of fullerene differing in sign of charge fullerenes. A significant part of the mass of negatively charged fullerenes spectra Fig 2 obtained optimum value of the discharge current is gradually at the decreases with increasing number of carbon atoms tail, which includes the structure and containing up to 150 and more carbon atoms. In this part of negatively charged ions fullerenes in intensity in 2 to 4 times higher than the background, but for ordinary fullerenes (\( C_{60}, C_{70} \ldots \)) the excess over the background of the peak of the fullerene has a value of 50-60.

MSs recorded from fullerite value obtained when the current discharge \( I = 80 \text{ A} \) are listed on Fig.3. The upper part (fullerenes with negative charge) has a lower intensity, than in Fig. 2 but run the marked features: intensity peak \( C_{84}^- \) greater, than the intensity \( C_{70}^- \), and in the spectrum of visible line \( C_{150}^- \).

**On the fullerenes with positive charge**

The lower part of Fig.3 shows the spectrum of positively charged fullerene. It’s clear that \( C_{84}^+ \) near the last observed, and the rest of the spectrum of the fullerene ions are indistinguishable from background structures. The calculation of data for Fig. 2 and an upper part of Fig.3 give an assumption that positively charged heavy fullerenes are neutralizing in moving from fullerite surface. This feature is a sign of any instability of positively charged fullerenes. Output calculations confirm the electrical parameters of fullerenes. According to the Table unlike the internal energy of the positively and negatively charged fullerene ratio (6) have more than 10 eV, and electron affinity \( (EA_i) \) have larger fullerenes with a large number carbon atoms. Parameter \( EA_i \) is a stabilizing factor structure of fullerene. Therefore, the mass spectrum on Fig. 2 and the upper part Fig.3, registered fullerenes \( C_{150}^+ \), but the spectrum of positively charged Fig. 3 ends near the \( C_{84}^+ \).

Comparison of the spectra shows that the mass spectrum of fullerene negatively charged are illustrating more details of the composition of analyte fullerite, and it has a greater stability of the electronic structure of fullerene ions with a negative charge. Therefore fullerene with a negative charge is preferably to use in electrodynamic processes.
REFERENCES


ISSN 2053-5783(Print), ISSN 2053-5791(online)
Fig.1. Projections of the parts of an arc discharge device with a hollow electrode and the conventional path A→B→C→D of fullerene formation [11, 19].

Table: The data of fullerenes [18], estimated value of electron affinity (EA_n) and the ionization energy (IE_n) of heavy fullerenes Fig. 2 [19].

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>C_20</td>
<td>0.693</td>
<td>10.04</td>
<td>10.733</td>
<td>-</td>
</tr>
<tr>
<td>C_60</td>
<td>2.67</td>
<td>7.57</td>
<td>10.24</td>
<td>2.667</td>
</tr>
<tr>
<td>C_70</td>
<td>2.87</td>
<td>7.41</td>
<td>10.28</td>
<td>2.75</td>
</tr>
<tr>
<td>C_106</td>
<td>3.33</td>
<td>7.06</td>
<td>10.39</td>
<td>3.41</td>
</tr>
<tr>
<td>C_112</td>
<td>3.39</td>
<td>6.98</td>
<td>10.37</td>
<td>-</td>
</tr>
<tr>
<td>C_122</td>
<td>3.47</td>
<td>6.91</td>
<td>10.38</td>
<td>-</td>
</tr>
<tr>
<td>C_146</td>
<td>3.64</td>
<td>6.78</td>
<td>10.42</td>
<td>-</td>
</tr>
<tr>
<td>C_150</td>
<td>3.66</td>
<td>6.76</td>
<td>10.42</td>
<td>-</td>
</tr>
</tbody>
</table>
Fig. 2. Mass fullerene arc hollow electrode, the discharge current $I = 70$ A, voltage $V \approx 20$ V, the parameters of the laser beam: $q = 2.6 \times 10^{-6}$ J, $f = 0.1$ s$^{-1}$; insertion: a- negative, b-positive[11, 19].
Fig. 3. Mass spectrum of the product obtained in ADHE-WGF with I = 80A after a double filtering; the parameters of the laser beam: q = 2.6x10^-6J, f = 0.1s^-1 [19].