Supplementary Materials
for

**Antioxidant Potential of Aqueous Dispersions
of Fullerenes C60, C70, and Gd@C82**by
Ivan V. Mikheev 1,\*, Madina M. Sozarukova 1,2, Dmitry Yu. Izmailov 3, Ivan E. Kareev 4, Elena V. Proskurnina 5, and Mikhail A. Proskurnin 1

1 Chemistry Department Analytical Chemistry Division of Lomonosov Moscow State University, Moscow 119991, Russia; mikheev.ivan@gmail.com

2 Kurnakov Institute of General and Inorganic Chemistry, Russian Academy of Sciences, Moscow 119991, Russia; s\_madinam@bk.ru

3 Faculty of Fundamental Medicine, Lomonosov Moscow State University, Moscow 119234, Russia; dizm@mail.ru

4 Institute of Problems of Chemical Physics of the Russian Academy of Sciences, 142432 Chernogolovka, Moscow Region, Russia; kareev@icp.ac.ru

5 Research Centre for Medical Genetics, Moscow 115522, Russia; proskurnina@gmail.com

\* Correspondence: mikheev.ivan@gmail.com; Tel.: +7(495)939-15-68 ext.101

[1 Aqueous fullerene dispersion preparation and material safety data sheet 3](#_Toc72677636)

[2 Quenching fullerene ability in aqueous media 5](#_Toc72677637)

[2.1 Quenching of ABAP-Luminol system in the presence of fullerene 5](#_Toc72677638)

[2.2 Quenching of Luminol system in the presence of fullerene 5](#_Toc72677639)

[**3** **Reference list** 8](#_Toc72677640)

# Aqueous fullerene dispersion preparation and material safety data sheet

Chemiluminescent systems can be influenced by the purity of the objects under study. Moreover, the action of the impurity component can be taken as a false positive mechanism. Thus, we carried out a deep purification of the AFDs. We have used the direct ultrasound-assisted preparation technique for AFDs of C60, C70, and Gd@C82 preparation [1] due to fewer impurity components and sedimental stability for more than 20 months.

Our experience with a sonication system for carbonaceous nanomaterials (hydrosols of nanodiamond) reveals titanium impurities [2] even for a short time (e.g., several minutes) of ultrasound treatment. Titanium accumulation rate was proportional to the prolonged ultrasound exposure time [3]. Our previous experiments proved that titanium and silicon are oxide species [4]. However, purification of AFD would not be resolved by now. The main idea of AFDs purification is primarily related to reducing the number of inorganic impurities (nanoparticles of titanium and silicon oxides), even considering the losses of the main component. Contamination of AFDs is associated with metal-carbon interactions [5]. It is essential to choose the correct filter system. We used commercially available syringe PVDF membrane filters of 0.22 or 0.45 μm. As for TiO2 NPs purification, a PVDF membrane can be used due to the high affinity of TiO2 for water [6]; the presence of hydroxyl moieties from TiO2 NPs contributed to improving the hydrophilicity, thereby improving the water ﬂux [7]. As shown in Table S1, filtration with a 0.22 or 0.45 μm PVDF filter completely removes TiO2 NPs fullerene dispersions.

Moreover, a 0.2 μm PTFE filter has low filtration efficiency toward titanium due to redox reactions between TiO2 NPs and Pu(III)/Pu(IV) at low pHs [3]. However, for practically neutral AFDs Table S1, the efficiency of PVDF filters is high. The presence of impurities can reduce or increase the effectiveness of antioxidants. The ability of TiO2 NPs to enhance the production of SAR and alter the antioxidant system in human osteoblast cells is shown [8]. Impurities may also adversely affect the reproducibility of results [9]. To evaluate the efficiency of fullerene antioxidant capacity, preparation of pure dispersion is an essential item.

As the AFD concentration after the combination of filters (prefilter 0.45 μm and primary filter 0.22 μm) is higher than after each filter alone (0.45 or 0.22 μm) (Table S1). Indeed, small pores (0.22 μm) clog faster, thereby reducing the filtration efficiency and, as a consequence, lowering the concentration of the target component [10]. Using prefiltration to increase filter life removes several significant limitations associated with losing fullerenes or endofullerenes. Prefiltration is widely used chromatography good laboratory practice [11].

As for SiO2 NPs purification, they remain in solution (approximately 65%, Table S1) after PVDF filters. SiO2 NPs pass through the membrane during ﬁltering with polytetraﬂuoroethylene (PTFE) membranes in a range of 0.1–1.0 μm of pore size [12]. In our opinion, we find a residual silica as typically silica SiO2 NPs can be formed with less than 220 nm in diameter [13]. Also, intermolecular and intramolecular interactions deal with probable silica surface with an adsorbed fullerene monolayer [14] and sorption of SiO2 NPs on the surface of fullerene clusters [15]. SiO2 NPs increased the capacity of the antioxidant enzymes superoxide dismutase (SOD), catalase (CAT), glutathione reductase (GR), and phenylalanine ammonia-lyase (PAL) in plants [16].

Thus, the total content of SiO2 NPs is less than the total fullerene content ratio in the stock solution by 10 to 170 times, which remains diluted. Therefore, it should not affect the measurements of antioxidant potential.

Table S1. Elemental composition of fullerene dispersions after preparation by direct ultrasound probe sonication (during 5h short-time duty) by inductively coupled plasma atomic emission spectroscopy (ICP-OES). Slurry sampling introduction in ICP-OES spectrometer. The uncertainty is ± 15%.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **Type of AFD or number of compounds** | **Syringe filter type** | ***c*Fullerene*,* ppm** | **pH\*** | ***c,*ppb** |
| **Ag** | **Al** | **As** | **B** | **Ba** | **Be** | **Cd** | **Co** | **Cr** | **Cu** | **Fe** | **Li** | **Mn** | **Mo** | **Ni** | **Pb** | **Si** | **Ti** | **V** | **Zn** |
| C60 | Pristine AFD (no filter using) | 1210 ± 50 | 7.35 | <5 | 290 | <5 | 320 | 17 | <0.1 | 25 | <1 | <1 | 80 | 75 | 16 | 45 | 273 | 60 | <1 | 2530 | 595 | 32 | 29 |
| 450 nm | 260 ± 20 | 7.44 | <5 | <10 | <5 | 105 | <1 | <0.1 | <0.5 | <1 | <1 | 7 | 7 | 2 | 5 | 10 | <1 | <1 | 1620 | <1 | <1 | 6 |
| 220 nm | 90 ± 10 | 7.40 | <5 | <10 | <5 | 105 | <1 | <0.1 | <0.5 | <1 | <1 | 7 | 7 | 2 | 4 | 10 | 4 | <1 | 1600 | <1 | 2.4 | 6 |
| Combination of 450 and 220 nm filters | 280 ± 24 | 7.45 | <5 | <10 | <5 | 105 | <1 | <0.1 | <0.5 | <1 | <1 | 7 | 7 | 2 | 4 | <5 | 3 | <1 | 1650 | 2.3 | <1 | 6 |
| C70 | 450 nm | 120 ± 10 | 7.55 | <5 | <10 | <5 | 90 | <1 | <0.1 | <0.5 | <1 | <1 | 5 | 5 | 2 | 2 | <5 | <1 | <1 | 1520 | <1 | <1 | <2 |
| Gd@C82 | 450 nm | 20 ± 4 | 7.74\*\* | <5 | <10 | <5 | 12 | <1 | <0.1 | <0.5 | <1 | <1 | 2 | 8 | 2 | 2 | <5 | <1 | <1 | 1450 | <1 | <1 | <2 |

\*, uncertainty was ± 0.10 pH units;

\*\*, pH of pristine AFD Gd@C82 was 7.89.

# Quenching fullerene ability in aqueous media

## Quenching of ABAP-Luminol system in the presence of fullerene

Interactions between the components of free-radical systems (ABAP, luminol) and aqueous fullerene dispersions (AFD) C60, C70, and Gd@C82 have been studied by fluorescent spectroscopy. Stern-Volmer approach (KSV) for quenching fluorescence analysis has been applied. Quenching occurs in static (with constant KS) or dynamic (with constant KD) interactions mechanism. To more speciation, fluorescence lifetime measurement should be applied. The magnitude of KS demonstrates that dynamic quenching cannot account for the decrease in intensity [17].

Normalization of fluorescence data at excitation 350 nm in Stern-Volmer coordinates is

$$\frac{I\_{0}}{I}=1+K\_{SV}[Q]$$

where $I\_{0}$ is the fluorescence intensity without a quencher; *I* is the fluorescence intensity with quencher addition; and Q is quencher concentration (M).

All known data are presented for 293K. Quenching of 1-pyrenencarboxilic acid by tris(malonate)-substituted C60 were KSV~4.4÷3.4)×104 M–1 [18]; for pristine C60 and fullerenol C60(OH)n with RNase A were 8.2×104 M–1 and 1.7×104 M–1 accordingly [19]; bovine (BSA) and human (HSA) serum albumins interaction between Gd@C82(OH)22 2.3 and 2.5×104 M–1 accordingly by a dynamic mechanism [20].

In this work, we conducted an estimation of quenching fullerene ability for AFD C60, C70, and Gd@C82 in the ABAP, luminol system. The magnitude of constant value was C60 ~ C70 > Gd@C82 (3.7 ± 0.1, 3.8 ± 0.1, and 2.9 ± 0.1)×104 M–1 which have good accordance with [18-20]. The fluorescence spectra are presented below (Figure S. 1). The different values of the Stern-Volmer constants for Gd@C82 are due to the different nature of the interactions [21]. It can be electrostatic interaction prevails and hydrogen bonds and van der Waals interactions [22] compared to non-endohedral fullerenes.

## Quenching of Luminol system in the presence of fullerene

In addition, we have estimated of quenching fullerene ability for AFD C60, C70, and Gd@C82 in only the luminol added system. The linearity of the Stern-Volmer plot indicates that only one quenching mechanism is possible; it could be either dynamic or static for more speciation lifetime measurements is needed. Luminol quenching by AFDs (Figure S. 2) have the magnitude of constant value in row C70 > C60 > Gd@C82 (7.4 ± 0.1, 5.7 ± 0.3, and 2.9 ± 0.1)×104 M–1. The values have good accordance with ABAP behavior, fullerene systems only for Gd@C82. Figure S. 1 and Figure S. 2 showed that AFDs could be quench both luminol and product of luminol triggered by ABAP radicals.

Figure S1. The fluorescent spectra of aqueous fullerene dispersions C60, C70, and Gd@C82 act as a quencher at 350 nm excitation wavelength, emission spectra registered 400–700 nm with maxima signal at 494 nm.

**Main areas:**

(1) Raman scattering peak of water at 397 nm;

(2) transparent blue area is luminol signal (cLuminol=2.0 μM);

(3) transparent green area is fluorescence of luminol triggered by ABAP radicals (cABAP=2.5 mM).

Registration conditions were excitation and emission slits 5 nm, scanning pitch 1 nm, integration time 0.1s, detector voltage 950 V, pH 7.4 (phosphate buffer) at 293K.

|  |  |
| --- | --- |
| (**a**) | Figure S2.The fluorescent spectra of luminol (cLuminol=2.0 μM) pristine and adding aqueous fullerene dispersions (AFD) C60, C70, and Gd@C82 act as a quencher at 350 nm excitation wavelength, emission spectra registered 400–700 nm with maxima signal at 428 nm.(**a**) AFD C60; (**b**) AFD C70; (**c**) AFD Gd@C82.Registration conditions were excitation and emission slits 5 nm, scanning pitch 1 nm, integration time 0.1s, detector voltage 950 V, pH 7.4 (phosphate buffer) at 293K. |
| (**b**) |
| (**c**) |

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