Synthesis, Spectroscopic Characterization and Photoactivity of Zr(IV) Phthalocyanines Functionalized with Aminobenzoic Acids and Their GO-Based Composites

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Abstract: Two complexes of bis(aminobenzoato)zirconium(IV) phthalocyanine and their graphite oxide-based composites were synthesized and characterized in respect of their photochemical properties. Structures of phthalocyanines were confirmed by Mass and infrared spectroscopies. The absorption and photoluminescence spectra were investigated to show various behavior of the complexes in different media (dimethyl sulfoxide and saline). Optical technique (monitoring variation of absorption spectra of diphenylisobenzofuran used as an indicator) was used to prove the generation of reactive oxygen species (ROS) by under light irradiation in the range of the first biological window. The photoactivity of the materials was compared and discussed in terms of their potential ability to be used in biomedical applications, for example, as photosensitizers in photodynamic therapy.

Keywords: graphite oxide (GO); phthalocyanine complexes; ROS generation; photoluminescence spectroscopy; photodynamic therapy

1. Introduction

Due to different abilities of phthalocyanines (Pcs), they have emerged huge interest in catalysis [1], semiconductors, photoconductors [2–4] and optical recording materials [5]. Despite their proficiency in the mentioned area, they are capable of use in medicine as photocytotoxic reagents [6]. Photodynamic therapy (PDT) is a clinically approved procedure for curing cancerous cells, which involves using photosensitizing reagents with the aid of an appropriate wavelength of visible or near infrared (NIR) light [7,8].

Reactive oxygen species (ROS), such as $^{1}\text{O}_2$, $\text{O}_2^{-*}$, $\text{OH}^{*}$ and $\text{H}_2\text{O}_2$, are responsible for cytotoxic effect which kills injured or microbial cells by a combinatorial effect of light [9,10]. Phthalocyanines are one of the most promising candidate in PDT; their role as photosensitizers and their activity in the red–NIR range (the first biological window) and high emission coefficients are not negligible [11–13]. Thus, their photochemical and photobiological properties have been widely reported in the literature [14–16].

Generally, Pcs and MPC complexes (M—Metal) are not soluble in water or dimethyl sulfoxide (DMSO). To use Pcs in pharmacology, good solubility of Pcs in water is preferred [17,18]. For this aim, metallophthalocyanines must be functionalized with hydrophilic organic/inorganic substituents.
in lateral position, or small inorganic counter-ions (Cl⁻ or OH⁻) or carboxylate groups should be substituted to metal cation of Pc complex in more or less axial position [17]. In our approach, the water solubility of the complexes is achieved by supplemental axial organic ligands attached to the central metal of the complex [19–21]. The choice of metal cation (Zr⁴⁺) was dictated by the fact that, in zirconium (or hafnium) phthalocyanine complexes, the coordination number of metal is usually 6. After the replacement of simple axial ligands, that is, Cl⁻ and OH⁻ by organic ligands, the coordination number of the metal increases to 8. Thus, when the complex is formed, metal has 2 additional coordination sites where at least 2 solvent molecules (water, DMSO, dimethylformamide (DMF), etc.) can be bonded. In addition, solubility may be significantly affected by hydrophilic groups present in axial ligands. These two factors cause that the zirconium complexes are relatively well soluble in water and DMSO [18].

Graphene family materials, such as graphene oxide and reduced graphene oxide, have been utilized in a large number of activities in biology field [22–24]. Recent studies showed that, for example, graphene oxide and its reduced forms have antibacterial properties towards Gram-positive, for example, *Staphylococcus aureus* and Gram-negative, for example, *Escherichia coli*, bacteria [25,26]. Antibacterial effect of graphene materials could be used to fabricate antimicrobial and biofilm resistant surfaces, for example, polymeric films for biomedical devices [27,28], antibacterial fabric materials [29] and graphene oxide-based membrane for water treatment [30–32]. There are different hypothesis for antibacterial effects of graphene materials. One is driven by spontaneous piercing of the membrane by nanoflakes of carboneous/graphene structures [33]. There is an interaction between basal hydrophobic region of graphene to hydrophobic inner region of plasma of membrane. Inducing cell damage in bacteria by extracting phospholipid molecules from the outer cell membrane [34] and direct oxidation of cellular components by graphene oxide [35] are the other pathways for integration of graphene oxide with bacterial cell membranes. Oxidative stress has another role to deactivate or kill bacteria which might be induced by generation of reactive oxygen species in the presence of graphene structures [35]. Embedding Pcs into graphene or similar nano supports might enhance their photo-physicochemical properties [36]. Macromolecules, such as Pc templates, have tendency to agglomerate in aqueous and biological media, which influence their absorption and emission properties. Due to a dye binding to a graphene carrier, agglomeration of molecules might be significantly reduced. Furthermore, transferring the dye into biological targets such as cells and tissues would be easier and less toxic to biological organisms while they are linked to a carrier [37]. In our previous research, new photosensitive nanomaterial with antibacterial photoactivity were introduced [38], where Pcs derivative was used together with graphene oxide and silver nanoparticles in a composite to enhance the antimicrobial effect.

In this paper we reported synthesis and photochemical characterization of two new zirconium(IV) phthalocyanine derivatives and their graphite oxide-based composites. The absorbance of ZrPcs with aminobenzoate ligands in dimethyl sulfoxide and saline were compared. Absorption spectroscopy was used to determine dependence of absorbance on Pc concentration and singlet oxygen generation under light irradiation for both Pcs and graphite oxide (GO) composites. Photoluminescent properties were also investigated.

2. Materials and Methods

2.1. Materials Syntheses

2.1.1. Pc Complexes

For axial substitution of zirconium (IV) phthalocyanine complex with 4-aminobenzoic acid (PABA) and 3,5-diaminobenzoic acid (DABA), a method described earlier was used [37,39]. Generally, in the first step, dichlorozirconium (IV) phthalocyanine was obtained by template synthesis from phthalimide (Alfa Aesar, Karlsruhe, Germany) and ZrCl₄ (ABCR, Karlsruhe, Germany) in 2-methylnaphthalene (Alfa Aesar) with heating. ZrPcI₂ was used as a precursor for a next step, where it reacted with 4-aminobenzoic acid (Alfa Aesar) or 3,5-diaminobenzoic acid (Alfa Aesar) in DMSO (Alfa Aesar) during
heating under reflux for about 6 h. The procedure is schematically shown in Figure 1. The complexes of bis (4-aminobenzoato) ZrPc and bis (3,5-diaminobenzoato) ZrPc were further abbreviated as bis(PABA)ZrPc and bis(DABA)ZrPc, respectively.

3. Results
3.1. Characterization of Pc Complexes and Composites

Two different zirconium (IV) phthalocyanines with mono and di-aminobenzoic functionalities axially substituted to the central metal ion and their GO composites were studied in this article (Figure 1). Graphite oxide flakes with dimensions of a few microns and a thickness of several dozen of carbon layers (TEM and SEM images of derived materials are presented in Figure S1 in the Supplementary Materials) and with different oxide groups were used to link functionalized phthalocyanines. To confirm the structures, FT IR and mass spectroscopy measurements were performed. Afterwards, the optical properties of these systems were studied and singlet oxygen generation was examined.

**Figure 1.** Synthesis of PcMCl₂ and bis (L) ZrPc complexes, where zirconium was used as metal ion (M) and 4-aminobenzoic acid (PABA) or 3,5-diaminobenzoic acid (DABA) were used as ligands (L).

2.1.2. Pc Composites

Highly oxidized graphite oxide (GO) was prepared according to the modified Brodie method [40] and composite materials were prepared according to a previous report [9]. GO powder (0.5 g) was suspended by ultrasonication (900 W, 22 kHz sonicator) in 5 mL of dimethylformamide (Alfa Aesar). Then, about 4 mg of bis (L) ZrPc, where L = PABA/DABA correspondingly, was added to the reaction vessel. Reaction mixture was placed in an ice bath and put on a magnetic stirrer. Dicyclohexylcarbodiimide (DCC, 0.06 g) (ABCR) was added to the mixture and the reaction was left in the ice bath on the magnetic stirrer for 12 h. After completion of the reaction, bis (L) ZrPc-modified GO materials were separated by centrifuge and washed 3 times with DMF and twice with warm ethanol to remove unreacted dye molecules. Composites were abbreviated as bis(PABA)ZrPc/GO and bis(DABA)ZrPc/GO consecutively.

2.2. Instrumentation

A spectrophotometer FLS980 (Edinburgh Instruments, Livingston, UK) with a xenon lamp was used for photoluminescence measurements. The spectra were recorded with the respective lamp and detector corrections. Absorption spectra of Pc solutions and Pc/GO suspensions were measured with a spectrophotometer Agilent CARY 5000 UV-Vis-NIR (Thermo Fisher Scientific, Walham, MA, USA). Polycrystalline IR spectra in the mid-IR region (400–4000 cm⁻¹) were measured using a Nicolet iS50
2.3. Materials Characterization

Structure of complexes were confirmed by ESI-MS analysis and IR spectroscopy. Mass spectra of compounds were recorded in DMF solutions. For FTIR spectra measurements samples were prepared as KBr pellets. For other optical measurements, stock solutions of ZrPcs in concentration of 0.5 mM were prepared by dissolving the compounds in dimethyl sulfoxide whereas the concentration of composites was about 18 mg/mL. Absorption spectra in the cases of two complexes of phthalocyanines, bis (PABA) ZrPc and bis (DABA) ZrPc, were recorded upon addition of 5 to 45 µL of ZrPc solution to the cuvette containing 2 mL DMSO and 5 to 125 µL of ZrPc to the cuvette containing 2 mL NaClaq (9 mg/mL) (the Beer-Lambert experiment, where absorbance versus dye concentration is plotted).

Generation of $^{1}O_2$ upon irradiation was shown by an indirect chemical method, using 1,3-diphenylisobenzofuran (DPBF) as a probe [41]. DPBF reacts with $^{1}O_2$ irreversibly, leading to a decrease in the absorption band at around 420 nm. The solution containing DPBF in DMSO (20 mg/mL) was prepared on the day of the analysis and stored in the dark. DPBF solution (14 µL) was added to a solution of the ZrPc complex (20 µL of solution in 2 mL DMSO) (to obtain absorbance level of a measured phthalocyanine solution in Q region around 1) or ZrPc composite (30 µL of suspension in 2 mL DMSO), which were then irradiated (from 10 s up to maximum 240 s) by red/NIR lamp (Philips, 100 W) which served as a light source that was located 30 cm far from a cuvette. Beer-Lambert graph for bis (PABA) ZrPc was plotted at 683 nm or 695 nm at which the absorption bands maxima were registered in DMSO and saline, respectively. For bis (DABA) ZrPc, graph data was registered at 690 nm and 700 nm in DMSO and saline, respectively. Photoluminescence and photoluminescence excitation spectra of phthalocyanine complexes were recorded in DMSO at $4 \times 10^{-5}$ M concentration.

3. Results

3.1. Characterization of Pc Complexes and Composites

Two different zirconium (IV) phthalocyanines with mono and di-aminobenzoic functionalities axially substituted to the central metal ion and their GO composites were studied in this article (Figure 1). Graphite oxide flakes with dimensions of a few microns and a thickness of several dozen of carbon layers (TEM and SEM images of derived materials are presented in Figure S1 in the Supplementary Materials) and with different oxide groups were used to link functionalized phthalocyanines. To confirm the structures, FTIR and mass spectroscopy measurements were performed. Afterwards, the optical properties of these systems were studied and singlet oxygen generation was examined.

FTIR spectra of bis (PABA) ZrPc and bis (DABA) ZrPc and their composites, bis (PABA) ZrPc/GO and bis (DABA) ZrPc/GO, display expected features (Figure 2), for example, bands at 3464, 3354 cm$^{-1}$ (Figure 2a) are attributed mainly to amine groups of ligands (Table S1, Supplementary Materials). The broad contour at 3500–3100 cm$^{-1}$ is related to OH groups on the GO surface (Figure 2b,d). The broad peaks observed in the region of 3500–3100 cm$^{-1}$ both in complexes and composites, are related to hydrogen bonding of NH$_2$ groups (Figure 2a,c). The narrower bands observed between 2841 and 3081 cm$^{-1}$ are assigned to stretching vibrations of CH groups. Strong bands in the 1516–1689 cm$^{-1}$ region are assigned mainly to stretching vibrations of carbonyl groups (C=O) coupled to vibrations of aromatic rings and bending vibrations of amine groups in bis (PABA) ZrPc and bis (DABA) ZrPc, respectively. Bending C–O vibrations are assigned to bands observed in the 1207–1288 cm$^{-1}$. Characteristic stretching vibrations of CNC groups are ascribed to bands observed in the 1207–1458 cm$^{-1}$ region, while their bending components contribute to many bands in the 505–736 cm$^{-1}$ and 843–904 cm$^{-1}$ regions. The bending in-plane vibrations of CH groups are related to many bands located between 946 and 1386 cm$^{-1}$, while out-of-plane aromatic CH are observed at lower wavenumbers (417–904 cm$^{-1}$).
The characteristic stretching vibrations of rings in Pc are assigned to bands between 1102 and 1600 cm\(^{-1}\). The in- and out-of-plane bending vibrations of rings contribute to bands observed in the 505–1088 cm\(^{-1}\) and 417–830 cm\(^{-1}\), respectively [42]. The characteristic bands of GO are clearly visible at 1068, 1612 and 1724 cm\(^{-1}\) for bis (PABA) ZrPc/GO and at 983, 1071, 1612 and 1725 cm\(^{-1}\) for bis (DABA) ZrPc/GO [43].

![Fourier-transform infrared (FTIR) spectra](image)

**Figure 2.** Fourier-transform infrared (FTIR) spectra of (a) bis (PABA) ZrPc, (b) bis (PABA) ZrPc/GO, (c) bis (DABA) ZrPc and (d) bis (DABA) ZrPc/GO.

By coupling reaction between phthalocyanine and GO, new linkages (amide bonds; Pc−NH−CO−GO) are formed in the system. Therefore, new peaks related to amide bonds are expected in the FTIR spectra of composites [44]. Three distinct differences can be found in FTIR spectra of composites that confirm the amide bond formation in the composites: (i) decrease in intensity of bands corresponding to NH\(_2\) stretching vibrations (Figure 2a,c); (ii) appearance of one narrow peak at 3325 cm\(^{-1}\) in bis (PABA) ZrPc/GO and 3326 cm\(^{-1}\) in bis (DABA) ZrPc/GO spectra (Figure 2b,d); (iii) and the increase of intensity and shifts of bands corresponding to the stretching vibrations of carbonyl (C=O) groups of amides at 1626 cm\(^{-1}\) in both composites (Figure 2a,c).

Mass spectroscopy analysis confirmed the structure of Pc complexes. In ESI-MS spectra of phthalocyanine complexes (Figure S2, Supplementary Materials), peaks with the molecular mass more than exact mass of complex could be seen due to one or two molecules of solvent (DMF) extracoordinated to metal atom of phthalocyanine complex as presented for bis (PABA) ZrPc in Figure 3. Coordinated solvent molecules are bonded more strongly with metal atom than axial ligand. Moreover, as it is shown in Figure 3, the structure breaks in ionization chamber of the spectrometer. Fragmentation starts with removal of amine functionalities from the ligand followed by detachment of phenone groups from the ZrPc structure. In this step, two molecules of DMF are bonded to the structure which will be removed later during ionization process. In the case of bis (DABA) ZrPc, the ESI-MS analysis give a molecular ion peak at \(m/z\)=908. In the case of bis (DABA) ZrPc bonded to one molecule of DMF (bis (DABA) ZrPc+DMF), peak at \(m/z\)=951 is observed. Signal at \(m/z\)=950 corresponds to the complex with one molecule of the solvent (DMF) while two amino groups are detached from the structure (bis (DABA) ZrPc+DMF-2NH\(_2\)). The signal at \(m/z\)=880 relates to the loss of two amino groups in bis (DABA) ZrPc complex. In ESI-MS, there is a possibility of formation of radical cations; especially easily oxidizable compounds, for example, hydrocarbon polyenes, polycyclic aromatic hydrocarbons, porphyrins and so forth. The ions observed by mass spectroscopy are quasimolecular ions created by the addition of a hydrogen cation \([M+H]^+\) or another cation, for example, Na denoted as \([M+Na]^+\) [45].
3.2. Spectroscopic Studies and Singlet Oxygen Generation

Optical properties of Pc complexes and composites were studied in two media (DMSO and saline—solution of sodium chloride in water). DMSO and saline were chosen for their compatibility in biological systems.

In the UV-Vis-NIR absorption spectra of MPcs, two specific regions are expected. Soret (or B) band usually is seen in the range between 300–400 nm. Band in the region between 620–750 nm named Q band is another characteristic feature. Both are seen in the spectra of the studied complexes. The intense absorption is visible at 345 nm (B band) and 685 nm (Q band) for both Pcs in DMSO (Figure 4a). Satellite peak is seen at 618 nm. Similar positions of the bands are seen for Pc complexes in saline—B band at 340 nm, Q band at 695 nm and satellite peak at 640 nm (Figure 4b). In transition metal complexes, these bands result from transition of electrons in molecular orbitals (d–d transitions). B band arise due to deeper π–π* transitions and Q band is attributed to the π–π* transition from HOMO to LUMO of Pc rings. Change of shape of the Q band and shift of band maxima (main and satellite) correspond with face-to-face dimerization of phthalocyanine complexes in water solution and conjugation of π-electron systems of two macrocycles.

Figure 3. Fragmentation pattern of phthalocyanine complexes according to ESI-MS analysis.

Figure 4. Absorption spectra of studied phthalocyanine complexes in (a) dimethyl sulfoxide (DMSO) and (b) saline. Molar concentration = 2 × 10^{-5} M.
The absorption spectra were also registered for different concentrations of complexes in DMSO or saline. The concentration of ZrPcs in DMSO was chosen to not exceed the value of Q band absorbance above 2.2 and the maximum concentration in saline used during experiment was about three times higher than in DMSO. The obtained spectra with the graphs showing the dependence of absorbance (at selected wavelength) on ZrPc concentrations are presented in Figure 5; Figure 6.

![Figure 5](image1.png)

**Figure 5.** Absorption spectra of different concentrations of bis(PABA)ZrPc in (a) DMSO (from 0.25 to 2.2 µM) and (b) saline (from 0.25 to 6.0 µM) and the results of Beer-Lambert linearity experiment (insets).

![Figure 6](image2.png)

**Figure 6.** Absorption spectra of different concentrations of bis(DABA)ZrPc in (a) DMSO (from 0.25 to 2.2 µM) and (b) saline (from 0.25 to 6.0 µM) and the results of Beer-Lambert linearity experiment (insets).

Capability of the complexes and composites as photosensitizers for singlet oxygen generation under irradiation was examined with 1,3-diphenylisobenzofuran which is a well-known reagent for this purpose. Absorption spectra of Pc complexes and composites in DMSO were recorded before and after addition of DPBF (Figures 7 and 8). Decrease of the DPBF absorbance (bleaching) ($\lambda_{\text{max}} = 418$ nm) in the presence of ZrPc complexes and composites was registered under different time of lamp irradiation.
Figure 7. Absorption spectra of (a) bis (PABA) ZrPc and (b) bis (DABA) ZrPc solutions in DMSO before and after addition of 1,3-diphenylisobenzofuran showing DPBF bleaching when exposed to light irradiation (due to singlet oxygen generation). (c) Irradiation time dependence of DPBF absorbance in the presence of ZrPcs.
Figure 8. Absorption spectra of (a) bis (PABA) ZrPc/GO and (b) bis (DABA) ZrPc/GO suspensions before and after addition of 1,3-diphenylisobenzofuran showing DPBF bleaching when exposed to light irradiation. (c) Irradiation time dependence of DPBF absorbance in the presence of ZrPc/GO composites.

Photoluminescence excitation and emission spectra of bis (PABA) ZrPc and bis (DABA) ZrPc in DMSO are shown in Figure 9. The photoluminescence excitation spectra were recorded monitoring the maximum of ZrPc emission band at 704 nm. Like in the case of absorption spectrum, two broad bands were observed: one in the range of 300–400 nm and the second one above 550 nm (here, only the satellite band around 620 nm was recorded). To avoid an overlap of emission band and excitation wavelength, the 620 nm excitation was used (instead of 690 nm where the maximum of Q band occurs). In the luminescence spectra, characteristic broad band in the range of 670–800 nm was observed. The measurement conditions did not allow to register spectra of complexes in saline nor composites.

Figure 9. (a) Photoluminescence excitation ($\lambda_{\text{em}} = 704$ nm) and (b) photoluminescence spectra ($\lambda_{\text{exc}} = 620$ nm) of bis (PABA) ZrPc and bis (DABA) ZrPc in DMSO.

4. Discussion

Dimerization, higher order association and self-aggregation of Pc rings usually take place in polar environments (polar solvents) that affects photochemical and photophysical properties of Pcs, for example, absorption intensity of complexes decrease during dimerization of metal phthalocyanine complexes. UV-Vis-NIR spectroscopy is a suitable mean to study this effect [46]. Shifts in Q band and changes in shape of this band indicate aggregation processes. In order to utilize Pcs in biomedicine, metal
phthalocyanines should have good solubility without aggregation in polar solvents [46]. Therefore, aggregation properties of the obtained complexes were studied at different concentration range (0.25–2.25 µM) in DMSO and (0.25–6.25 µM) in saline.

First of all, the registered absorption spectra of the complexes are common for metal phthalocyanines in non-polar (DMSO) and polar (saline) solvents showing different width and relative intensities of the bands. Shape of the spectra shown in Figures 5b and 6b resulted from dimerization of Pc structures in aqueous solution. Looking at the spectra registered for various ZrPc concentrations, the findings were in agreement with Beer-Lambert law linearity experiment in DMSO showing that there is no clear evidence of molecules aggregation (dimerization and further agglomeration) in this solvent for the studied concentration range (insets in Figures 5a and 6a). The same should be for other organic solvents as verified in our earlier works [20,21]. In case of using saline solution for both complexes, deviation from Beer-Lambert linearity law was observed for ZrPc concentration higher than 4 and 3 µM, for bis (PABA) ZrPc and bis (DABA) ZrPc, respectively (insets in Figures 5b and 6b). Non-linear dependence indicated clearly high degree of molecules aggregation. Lower concentration limit for bis (DABA) ZrPc may be observed due to higher polarity of DABA ligands (higher tendency to dimerization than in the case of PABA) [20]. Eventually, in the spectra of GO-based structures functionalized with ZrPc derivatives, weak bands at around 690 nm were seen (Figure 8a,b) confirming the presence of phthalocyanines in the composites.

Photoluminescence excitation spectra (Figure 9) showed that two ranges of light wavelengths can be used for samples excitations, that is, UV/blue and red/NIR. Red (here 620 nm) excitation confirmed the presence of red/NIR luminescence of ZrPc complexes. Photoluminescence was not recorded for complexes in saline or for composites. In the first case, the aggregation of ZrPc might be the reason for low luminescence properties. In the case of composites, low content of ZrPc on GO flakes (as indicated by the absorption spectra) would explain lack of seen light emission.

Broadband red/NIR light emitting lamp was used for singlet oxygen generation tests. This commercially available light source was suitable for ZrPcs photoactivation. Both phthalocyanine complexes produced reactive oxygen species with similar effectiveness (Figure 7). The DPBF absorption band was not seen after 2 min of irradiation.

Despite the fact that the concentration of PCs was much lower in the composites than in the solutions, singlet oxygen generation was still observed for these samples but the amount of reactive species was not enough high to react with the DPBF molecules after 4 min of irradiation. Nevertheless, the idea to link chosen PCs to a GO carrier and to maintain their photoactivities was proved to be correct. The results shown in Figure 8c marked that the activity of bis (DABA) ZrPc/GO is slightly higher (faster DPBF bleaching). Differences in the amount of \(^{1}\text{O}_2\) generated by bis (L) ZrPcs may depend on their structure. Bis (PABA) ZrPc with two amino groups of two ligands is geometrically favorable (less hindrance for para position) but bis (DABA) ZrPc, with four amino groups in meta position, provides better connection/linkage to graphite oxide as support (mainly due to the possibility to form more hydrogen bonding but also through \(\pi-\pi\) stacking or covalent bonding to carrier surface). To increase the photoactivity of composites, one could think of loading more of Pc complexes on the GO structure. In this study, synthesis of composites was already performed in a way to ensure relatively high concentration of ZrPc and it would be difficult to obtain higher phthalocyanine content. But it must be noted that structure of complexes with different functionalities absolutely affects the attachment process that further influence on the \(^{1}\text{O}_2\) generation as reported in our previous studies of other ligands bonded to ZrPc [38,47].

In conclusion, the generation of singlet oxygen has been a subject of great interest in different disciplines, from polymer science to medicine. High chemical reactivity of singlet oxygen can be utilized in biological systems, for example, in PDT [48] including antiseptic effect. Photoactivity capacity of the studied materials (both complexes and composites) was proved by the test of singlet oxygen generation using DPBF. Comparison between complexes and composites showed better \(^{1}\text{O}_2\) generation in ZrPcs. From excitation and emission spectra of Pc complexes, one could conclude that
these phthalocyanine compounds are good candidates for PDT photosensitizers, since excitation at 600–700 nm and emission between 700–800 nm are essential to penetrate deep enough into tissues [38].

**Supplementary Materials:** The following are available online at http://www.mdpi.com/2311-5629/6/1/1/s1, Figure S1: TEM and SEM images of graphite oxide flakes with zirconium(IV) phthalocyanine derivatives; Figure S2: Mass spectra of (a) bis (PABA) ZrPc, (b) bis (DAPA) ZrPc; Table S1: IR wavenumbers and relative intensities of observed bands together with proposed assignment for the studied complexes and composites.

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