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Porphyran-based bridged silsesquioxane nanoparticles for targeted two-photon photodynamic therapy of zebrafish xenografted with human tumor

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Abstract

Background: Bridged silsesquioxane nanoparticles (BSNs) recently described represent a new class of nanoparticles exhibiting versatile applications and particularly a strong potential for nanomedicine.

Aims: In this work, we describe the synthesis of BSNs from an octasilylated functional porphyrin precursor (PORBSNs) efficiently obtained through a click reaction. These innovative and very small-sized nanoparticles were functionalized with PEG and mannose (PORBSNs-mannose) in order to target breast tumors in vivo.

Methods and Results: The structure of these nanoparticles is constituted of porphyrins J aggregates that allow two-photon spatiotemporal excitation of the nanoparticles. The therapeutic potential of such photoactivable nanoparticles was first studied in vitro, in human breast cancer cells in culture and then in vivo on zebrafish embryos bearing human tumors. These animal models were intravenously injected with 5 nL of a solution containing PORBSNs-mannose. An hour and half after the injection of photoactivable and targeted nanoparticles, the tumor areas were excited for few seconds with a two-photon beam induced focused laser. We observed strong tumor size decrease, with the involvement of apoptosis pathway activation.

Conclusion: We demonstrated the high targeting, imaging, and therapeutic potential of PORBSNs-mannose injected in the blood stream of zebrafish xenografted with human tumors.

KEYWORDS
bridged silsesquioxane nanoparticles, human tumor targeting, photodynamic therapy, two-photon excitation
1 | INTRODUCTION

Nanotechnology represents a major advance in the field of medicine and has started to play an important role in the development of anticancer therapeutics.\textsuperscript{1,2} In the course of the development of more efficient, biocompatible, multifunctional, and stealth nanoparticles for theranostics, we turned to bridged silsesquioxane nanoparticles (BSNs).\textsuperscript{3} Indeed, BSNs were recently described and represent a new class of nanoparticles exhibiting versatile applications in different areas of research such as photovoltaics, electrochromism, heterogeneous catalysis, environment, and nanomedicine.\textsuperscript{4} They are obtained through the sol-gel condensation in mild conditions, using organic precursors possessing at least two trialkoxysilane groups without the presence of any silica source. Despite their diverse properties and various relevant applications, several features account for their interest for biomedical use. In fact, the size, morphology, and surface are controllable; the high organic content of BSNs (20-80 wt%) can be used for theranostic purposes with a strong potential and a good biocompatibility.\textsuperscript{5} We demonstrated in a previous work that these BSNs mainly composed of photosensitizers were very effective for photodynamic therapy (PDT) of cancer cells under two-photon excitation (TPE).\textsuperscript{5}

TPE is very attractive for nanomedicine applications\textsuperscript{6,7} as it allows a deep penetration of the near-infrared beam down to 2 cm in tissues\textsuperscript{8} and a high spatiotemporal resolution presenting a strong interest for imaging and cancer detection. Importantly, TPE-PDT has demonstrated a high potential for cancer therapy\textsuperscript{9-26} and particularly for small-sized tumor treatment.

Today, the evolution of nanotechnology, involving or not PDT, requires high competitive and accurate methodologies to determine the therapeutic effect and the biocompatibility of the new nanostructured materials for biomedical applications. In particular, the elaboration of integrated models for in vivo studies of nanomaterials becomes a priority for the research in the field of nanomedicine. In 2010, the European directive on the protection of laboratory animals was revised, and it was requested to find alternative methods to the use of sensitive animals such as small mammals. The embryo is not considered a sensitive animal, and its use is privileged. The zebrafish embryo is a very popular animal model because it is easy to breed, with a fast development, and possesses 80\% of gene homology with humans.\textsuperscript{27} Previous studies have shown that it is a powerful model organism for the study of human biology, being well suited to both developmental and genetic analyses.\textsuperscript{28} Importantly, they can easily develop human tumors due to their lack of immune system in this embryonic state.\textsuperscript{29} Moreover, the embryos that develop outside the mother are transparent, which is a great advantage in our case for tumor imaging. For all these crucial benefits, we propose to use this integrated animal model for investigations on innovative and targeted BSNs for anticancer properties under TPE. Recently, we described new porous porphyrin-based organosilica nanoparticles of a 250-nm average diameter, which were endocytosed in MDA-MB-231 breast cancer cells before being injected in zebrafish embryos.\textsuperscript{30} After 2 days, the tumor developed, and TPE-PDT was successfully performed on the xenografted tumor, showing the proof of concept of TPE-PDT with zebrafish embryos. However, this experiment could be related to an intratumoral injection of nanoparticles, and the size of the nanoparticles which did not allow us to inject in the tail vein was a limit in this work. This is the reason why we decided to synthesize small organic nanoparticles which was very challenging.

Herein, we describe the preparation of small-sized non-porous porphyrin-based BSNs (PORBSNs) from an octasilylated porphyrin precursor, anchored with both stealth polyethylene glycol (PEG) groups to bring hydrosolubility and mannose groups in order to target breast cancer cells through interactions with lectins overexpressed in MDA-MB-231 breast cancer cells.\textsuperscript{31} The therapeutic potential of these small-sized PORBSNs-mannose is presented in vitro using human breast adenocarcinoma cells (MDA-MB-231) and in vivo through intravenous injection of the nanoparticles in zebrafish xenografted with this cancer cell line. The biological study of TPE-PDT demonstrated their strong efficiency after only a few seconds of irradiation on cell cultures and on human tumors in the zebrafish model.

2 | MATERIAL AND METHODS

2.1 | Materials

Cetyltrimethylammonium bromide (CTAB, 99%), sodium hydroxide (97%) DMSO, and AcOH were purchased from Sigma-Aldrich. EtOH was purchased form Carlo Erba. Aminoundecyltrimethoxysilane and (6-[2-(2-[2-methoxy-ethoxy]-ethoxy)-ethoxy]-hexyl)trimethoxysilane were purchased from SIKEMIA. p-[N-(2-Ethoxy-3,4-dioxocyclobut-1-enyl)amino]phenyl-α-D-mannopyranoside was synthesized as described.\textsuperscript{32}

2.2 | Analytical techniques

UV-vis absorption spectra were recorded on a Hewlett-Packard 8453 spectrophotometer using correction factors supplied by the manufacturer. N\textsubscript{2} adsorption isotherms were measured using a TRISTAR 3000 gas adsorption analyzer instrument, and the specific surface area was determined using the BET method. TEM analysis was performed on a JEOL 1200 EXII instrument. Dynamic light scattering analyses were performed using a Cordouan Technologies DL 135 Particle size analyzer instrument.

2.3 | Synthesis of porphyrin-based bridged-silsesquioxane nanoparticles (PORBSNs)

A mixture of CTAB (250 mg, 0.78 mmol), distilled water (120 mL), and NH\textsubscript{4}OH (0.2 M aqueous solution, 1.2 mL) was stirred at 80°C for 2 hours at 750 rpm in a 250-mL three-neck round bottom flask. Then, the octasilylated porphyrin\textsuperscript{30} (100 mg, 0.0327 mmol, in 2 mL of absolute ethanol) was added, and the mixture was stirred for
30 hours at 80°C. Afterwards, the solution was cooled to room temperature while stirring and stored in the fridge.

### 2.4 Synthesis of mannose-functionalized porphyrin-based bridged-silsesquioxane nanoparticles (PORBSNs-mannose)

Twenty milligrams of aminoundecytrimethoxysilane (0.09 mmol) was dissolved in 10 mL of anhydrous EtOH; 349 mg of p-[N-(2-ethoxy-3,4-dioxocyclobut-1-etyl)amino]phenyl-α-D-mannopyranoside (squalate mannose) (0.88 mmol; 10 eq.) was added to the solution. The mixture was stirred at 50°C overnight and then cooled to room temperature. The crude mixture was centrifuged (15 000 rpm; 5 min). The supernatant was removed, and the pellet was resuspended in EtOH. This operation was renewed once. After removal of the supernatant, the product was recovered as a pale yellow paste. This paste was stored without further drying.

Twelve milligrams of this paste was dissolved in 1.3 mL of DMSO, and 100 mg of 6-{β-[2-[2-methoxy-ethoxy]-(ethoxy)-hexyl]trimethoxysilane was added. This solution was added to 8 mL of the solution of PORBSNs and the mixture stirred at 700 rpm and 50°C for one night. After reaction, excess reagents and CTAB were removed by dialysis (cutoff 12 kDa) in a mixture of EtOH/H2O/AcOH: 1/1/0.07 for 24 hours. The operation was repeated twice. The suspension was then dialyzed in EtOH for 3 × 24 hours and stored at room temperature in EtOH.

### 2.5 Cell culture

Human breast adenocarcinoma cells MDA-MB-231 expressing (or not) green fluorescent protein (GFP) (purchased from ATCC) were cultured in DMEM Media–GlutaMAX-I (containing 4.5 g.L⁻¹ of D-glucose) supplemented with 10% fetal bovine serum and 50 μg mL⁻¹ gentamycin and allowed to grow in a humidified atmosphere at 37°C under 5% CO₂.

### 2.6 Cytotoxicity measurement

For in vitro cytotoxicity analysis, MDA-MB-231 cells were seeded into a 96-well plate, 1000 cells per well in 200 μL of culture medium, and allowed to grow for 24 hours. Then, cells were treated with increasing concentrations of PORBSNs-mannose, and after 3 days, a MTT assay was performed as previously described.

### 2.7 OPE or TPE-PDT experiments

For in vitro OPE-PDT, MDA-MB-231 cells were seeded into a 384 multiwell glass-bottomed plate (thickness 0.17 mm), with a black polystyrene frame, 500 cells per well in 50 μL of culture medium, and allowed to grow for 24 hours. Then, cells were treated with 80 μg mL⁻¹ PORBSNs-mannose, and 24 hours after, cells were submitted (or not) to laser irradiation, with the Carl Zeiss Microscope (laser power input 3 W). Half of the well was irradiated at 800 nm by three scans of 1.57-seconds duration in four different areas of the well. The laser beam was focused by a microscope objective lens (Carl Zeiss 10×/0.3 EC Plan-Neofluar). The scan size does not allow irradiating more areas without overlapping. After 2 days, the MTT assay was performed as previously described and was corrected according to the following formula: Abs control – 2 × (Abs control × Abs PORBSNs-mannose).

### 2.8 ROS imaging under one or two-photon excitation

The detection of intracellular reactive oxygen production (ROS) was realized using DCF-DA Cellular ROS Detection Assay Kit (abcam). For ROS measurement under monophotonic excitation, MDA-MB-231 cells were seeded in a 12-well plate and treated with PORBSNs-mannose at 80 μg mL⁻¹. After 24 hours, cells were rinsed and incubated 45 minutes at 37°C with 2,7-dichlorofluorescein diacetate (DCFDA) at 20 μM. Then, cells were rinsed and submitted or not to an irradiation with a mercury lamp, during 10 minutes (λ = 420-440 nm). Green luminescence translates the generation of ROS detected at 535 nm.

For ROS measurement under biphotonic excitation, cells were seeded in bottom glass dishes (World Precision Instrument, Stevenage, UK) at a density of 10⁶ cells·cm⁻² and submitted to the same protocol with an incubation with a LSM 780 LIVE confocal microscope (Carl Zeiss, Le Pecq, France), at 800 nm (3 × 1.57 seconds) followed by washing and imaging at 535 nm.

### 2.9 Two-photon fluorescence imaging

The day prior to the experiment, MDA-MB-231 cells expressing GFP were seeded onto bottom glass dishes (World Precision Instrument, Stevenage, UK) at density of 10⁶ cells cm⁻² and submitted to the same protocol with an incubation with a LSM 780 LIVE confocal microscope (Carl Zeiss, Le Pecq, France), at 800 nm for PORBSNs-mannose, 488 nm for GFP and 561 nm for cell membranes. All images were performed with a high magnification (63×/1.4 OIL DIC Plan-Apo).
2.10 | Cell preparation for injection

The day of the experiment, the cells were washed twice, trypsined, and then stained with 2 μg·mL⁻¹ of Dil Stain (1.1'-dioctadecyl-3,3',3'-tetramethylindocarbocyanine perchlorate ("DiI"; DilC18(3))) (Invitrogen), 15 minutes at 37°C.

In parallel, the cells were deposited on a counting chamber (MARIENFELD SUPERIOR), to determine the number of cells per volume unit of liquid. The stained cells were diluted to final concentration of 10⁸ cells·mL⁻¹ in Dulbecco's phosphate-buffered saline (DPBS) supplemented with 10% fetal bovine serum. The cells are maintained in ice until injection, which must be done within 3 hours after the preparation of the cells.

2.11 | Zebrafish used model and cancer cell injection

Zebrafish experiments were done according to the guidelines of the European Community Council directive 2010/63/EU and to the accreditation number of the zebrafish facility A34-172-37 of the University of Montpellier. Casper line, zebrafish pigmentation mutants, were purchased from the Zebrafish International Resource Center (ZIRC) as embryos and raised to adulthood in Roussel lab's facilities. Only fish directly from ZIRC or their F1 offspring were used as egg producers to avoid inbreeding effects. Embryos were obtained from pairs of adult fish by natural spawning and raised at 28.5°C in tank water. Embryos and larvae were staged according to Kimmel et al.⁵³

Zebrafish larvae were manually dechorionated and anesthetized. One hundred cells per nanoliter were injected into the perivitelline space on embryos/early larvae aged 24 to 30 hours postfertilization (hpf); between 5 and 10 nL was injected each time. The injected or not larvae were allowed to develop at 32°C. It was observed at 24 hours (hpi) under an Olympus MVX10 epifluorescence microscope. Only the larvae with homogeneous and comparable xenografts were kept.

PORBSNs-mannose are diluted to 4 mg mL⁻¹ in water containing 1% phenol red and then intravenously injected (5 nL) on xenografted zebrafish larvae of 4 days postfertilization.

For each condition, the injected (or not) embryos were divided into two sets: one that will be irradiated and the other not. In the order to follow and compare the xenografts evolution, the irradiated or not larvae were allowed to develop at 32°C until the sixth day.

2.12 | Two-photon excited photodynamic therapy of zebrafish

For two-photon excited irradiation, zebrafish larvae were placed into a multiwell glass bottom with the Carl Zeiss Microscope (laser power input 3 W). The laser beam was focused by a microscope objective lens (Carl Zeiss 10×/0.3 EC Plan-Neofluar). The tumor area of zebrafish larvae was irradiated at 800 nm by three scans of 1.57-second duration.

2.13 | Zebrafish fluorescence imaging

The injected or not larvae were imaged before and 2 days after two-photon irradiation. Each larva was placed into a multiwell glass bottom (thickness 0.17 mm) and scanned with a LSM 780 LIVE confocal microscope (Carl Zeiss, Le Pecq, France), at 561 nm. All images were performed by a confocal microscope objective 20× (Plan-APOCHROMAT 20×/0.8 M27). The quantification of fluorescence intensities and tumor volumes in zebrafish embryos were performed by ImageJ software after 3D reconstruction of Z-stacks. For reconstitution of tumor treated with PORBSNs-mannose and laser, 47 Z-stacks were performed with a slice thickness of 2.72 μm each one. For reconstitution of tumor treated with PORBSNs-mannose alone, 274 Z-stacks were necessary with a slice thickness of 3.86 μm each one. For finish, reconstitution of tumor submitted to laser only, 242 Z-stacks were performed with a slice thickness of 4.4 μm each one.

2.14 | Immunofluorescence staining in toto zebrafish

The embryos were fixed at 6 days postfertilization in paraformaldehyde (PFA) 4% in PBS + 0.5% triton at room temperature (RT) for 3 hours, under hood, and then washed with PBS at least 3 times. Fixed embryos were permeabilized using Triton % in PBS for 2 hours, at RT. In order to saturate the potential unspecific sites antibodies, the embryos were then blocked with blocking solution, containing: PBS 1X + BSA 1X + DMSO 1X + 10% Donkey serum (Sigma-Aldrich D9663) for 1 day at 4°C.

Primary cleaved caspase-3 (Asp175) antibodies (Cell Signaling Technology) were used for specific labeling of the apoptosis signal. The recommended antibody dilution is 1:400 in a blocking solution containing 2% of Donkey serum. The larvae were incubated over the weekend at 4°C and then washed during the day in PBS. At the end of the day, the larvae were labeled with the secondary antibody conjugated to Cy5 at the dilution of 1:500 and incubated overnight at 4°C. Then, the larvae were washed whole day in PBS and stained with Hoechst dye at the last wash at the concentration of 7 μg mL⁻¹ during 40 minutes and finally washed, again, at least three times with PBS. The immunostained larvae were mounted flat onto transparent slides with a coverslip (using Dako fluorescent mounting medium).

2.15 | Zebrafish 3D imaging

Confocal fluorescence microscopy was performed using a Leica DM2500CSQ upright microscope with a Leica TCS SPE confocal scan head, differential interference contrast (DIC) optics, and a SuperZGalvo SPE z-step controller. The acquisition of three-dimensional stacks was performed with 63 × Leica Apo oil 1.15 NA. Final image analysis and reconstitution were performed using Imaris X64 8.02 software.
3 | RESULTS

3.1 | PORBSNs and PORBSNs-mannose: Synthesis and characterization

The synthesis of PORBSNs was performed from a porphyrin precursor possessing eight triethoxysilyl groups (Scheme 1). The sol-gel reaction was carried out in diluted conditions using cetyltrimethylammonium bromide (CTAB) as a template and NH₃ as a catalyst. The condensation was performed for 30 hours at 80°C yielding a colloidal suspension of small-sized PORBSNs. After cooling, the suspension was stable and was stored in the fridge without purification in order to keep the colloidal stability.

Nitrogen sorption performed on the dried nanoparticles revealed a low specific surface area (20 m² g⁻¹) indicating that the nanoparticles were not porous. This was confirmed by transmission electron microscopy (TEM) (Figure 1A) that depicts mainly nanoparticles of small size (between 20 and 30 nm), while some bigger nanoparticles (100 nm) were also observed. These nanoparticles were much smaller than our previously reported BS nanoparticles obtained from a tetrasiylated porphyrin precursor, which showed a size distribution centered at 106 nm. Dynamic light scattering (DLS) of the suspension of PORBSNs (Figure 1B) suggests that small-sized nanoparticles prevailed in a solution with a distribution centered at 20 nm. The biggest nanoparticles (50-100 nm) represented a minor contribution. PORBSNs were then functionalized with PEG and mannose following our recently published method for small-sized mesoporous silica nanoparticles (Scheme 1). Well-dispersed small-sized nanoparticles were obtained as suggested by TEM (Figure 1C) and DLS (Figure 1D), with a decrease of the mean hydrodynamic diameter to 7 nm with a narrow size distribution. This can be explained by the elimination of excess CTAB during the procedure of grafting, due to extensive dialysis with AcOH/ETOH which led to disaggregation of the nanoparticles. PORBSNs-mannose were stored in ETOH at 0°C, and the suspension was stable at concentrations as high as 14 mg mL⁻¹.

UV-vis spectra of the porphyrin precursor, PORBSNs, and PORBSNs-mannose are presented in Figure 2.

The metallated porphyrin showed the characteristic Soret band at 425 nm and the two Q bands at 558 and 600 nm. PORBSNs showed an enlarged Soret band at 440 nm and the two Q bands at 564 and 604 nm. Compared with the precursor, the bands were redshifted in PORBSNs in agreement with the formation of J aggregates of porphyrins, which endows the nanoparticles with two-photon sensitivity. PORBSNs-mannose showed the characteristic bands of the J porphyrin aggregates and of the phenyl-squarate moiety at 320 nm, confirming a successful functionalization of BSNs with mannose.

3.2 | In vitro biological study

The biocompatibility of PORBSNs-mannose was first investigated on human cancer cells in culture. The absence of toxicity of PORBSNs-mannose in the dark was verified on human breast adenocarcinoma cells (MDA-MB-231) incubated 72 hours with increasing concentrations of PORBSNs-mannose. At the end of the experiment, the living cells were quantified by a colorimetric assay (MTT). The experiment is described in Figure 3 and showed no significant cell death in a range of nanoparticles from 10 to 200 μg mL⁻¹.

The one-photon excited PDT (OPE-PDT) efficiency of PORBSNs-mannose was first investigated on MDA-MB-231 breast cancer cells. We studied OPE-PDT and the reactive oxygen species (ROS) production of the nanoparticles with a monophotonic excitation (420-440 nm for 10 minutes, using the mercury lamp of a standard fluorescence microscope). MDA-MB-231 cells incubated for 24 hours with 80 μg mL⁻¹ PORBSNs-mannose underwent a significant cell death of 44% after 10 minutes of irradiation (Figure 4A). The PDT effect is due to the production of ROS which leads to the destabilization of cell and organelle membranes, DNA alteration, and finally cell death. We verified that the phototoxicity observed after nanoparticles treatment and irradiation was due to ROS production. For this, cells were treated with PORBSNs-mannose in the same conditions as those used for PDT, but before irradiation, 20-μM 2,7-dichlorofluorescein diacetate (DCFDA) were added in a culture medium (Figure 4B). The green luminescence detected at 535 nm was characteristic of the generation of ROS.

We then investigated two-photon excited PDT (TPE-PDT) with PORBSNs-mannose. As shown in Figure 5A, PORBSNs-mannose led to 50% cancer cell death after a few seconds of excitation.
(3 × 1.57 seconds at 800 nm) with a femtosecond pulsed laser. This effect was due to the production of ROS in the cells treated with PORBSNs-mannose and irradiated with TPE, as demonstrated in Figure 5B.

The internalization of the nanoparticles in cancer cells was then assessed after 24 hours of incubation at 80 μg mL⁻¹ with a two-photon Carl Zeiss confocal microscope at low power (3%, input power 3 W, excitation at 800 nm) (Figure 6). Bright fluorescent dots were observed with PORBSNs-mannose and were localized in the cells, thus demonstrating nanoparticle uptake.

To investigate in vivo TPE-PDT, we used zebrafish larvae as an integrated model for human tumor. The efficiency of PORBSNs-mannose with this model was studied. First, MDA-MB-231 human breast adenocarcinoma cells (stained with red Dil stain) were implanted in the perivitelline space of embryos 24 to 30 hpf. The successfully injected embryos were sorted using the red luminescence of MDA-MB-231. The homogeneous xenografted zebrafish embryos were then kept for 2 days until the development of the tumors. Four days postfertilization (dpf), the embryos were intravenously injected or not with PORBSNs-mannose and then submitted or not to TPE irradiation 1.5 hours after the injection of the nanoparticles. Finally, the zebrafish embryos were bred 2 days longer to evaluate the evolution of the xenografts.
FIGURE 4  Phototoxicity study of PORBSNs-mannose on MDA-MB-231 cells treated with 80 μg mL\(^{-1}\) of nanoparticles. A, PDT efficiency of PORBSNs-mannose after cell irradiation at 420 to 440 nm for 10 min. Cell death quantification was realized by MTT assay 48 h after irradiation. Data are mean ± standard deviation of three experiments. *\(P \leq 0.05\) statistically different from control (Student t test). B, ROS production by MDA-MB-231 cells after PORBSNs-mannose incubation. Cells are incubated with DCF-DA and irradiated (420-440 nm, 10 min). Green luminescence translates ROS production by cancer cells (scale bar 50 μm).

FIGURE 5  In vitro TPE-PDT experiment and ROS production of PORBSNs-mannose incubated for 24 h at 80 μg mL\(^{-1}\) with MDA-MB-231. A, Cell survival measurements are achieved by MTS assay 48 h after irradiation (at 800 nm with a pulsed laser for 3 × 1.57 s). Data are mean values ± standard deviation from three independent experiments. *\(P \leq 0.05\) statistically different between laser OFF and laser ON (Student t test). B, Reactive oxygen species (ROS) were detected on MDA-MB-231 cancer cells incubated or not with PORBSNs-mannose (80 μg mL\(^{-1}\) for 24 h) and added with DCF-DA 45 min before irradiation (at 800 nm with a pulsed laser for 3 × 1.57 s). Generated ROS were monitored by the green fluorescence of 2,7-dichlorofluorescein at 535 nm (scale bar 50 μm).
As shown in Figure 7A, a clear evolution of the xenografts was observed 6 dpf with live larvae injected with MDA-MB-231 cells and PORBSNs-mannose without TPE. Then, TPE was performed on larvae injected with MDA-MB-231 cells without previous PORBSNs-mannose treatment, in order to demonstrate that the irradiation at 800 nm did not affect the zebrafish embryo survival or tumor development. In contrast, the intravenous injection of PORBSNs-mannose followed by the two-photon irradiation significantly reduced the xenografts size (Figure 7A). This experiment was repeated and finally done four times. As reported in Figure 7B, in all cases, the xenografted animals showed an increase in tumor size while when zebrafish embryos bearing tumor were also injected with PORBSNs-mannose and irradiated at the tumor area during 3 × 1.57 seconds, a strong disappearance of the tumor could be observed, and abundant white dots were revealed in the presence of anti-caspase 3 antibodies, thus specifying apoptotic death of cancer cells (Figure 8F).

**DISCUSSION**

In this work, we reported small-sized PORBSNs-mannose constituted of porphyrins and anchored with mannose moieties. These nanoparticles present a high potential in biomedical applications. First, the use of BSN allows a good biocompatibility because of the high quantity of organic moieties in this hybrid nanomaterial. Secondly, the porphyrins organized in J aggregates inside the BSN could be excited with a pulsed laser in near-infrared wavelengths. This means that, in these nanoparticles, besides exciting by a monophotonic source of laser, porphyrins were excited with a two-photon laser with a strong efficiency. Currently, the two-photon laser excitation in the near infrared region leads to an increased penetration depth in tissues with less photodamages and with a spatiotemporal resolution suitable for small-sized tumors. Thirdly, the anchoring of PEG and mannose moieties at the surface of PORBSNs-mannose demonstrated biocompatibility and effectiveness for cancer therapy.

The in vitro experiments we performed on human breast cancer cells in culture have confirmed that PORBSNs-mannose were safe.
and induced cancer cell death upon irradiation with OPE or TPE. In a second time, we developed an animal model to study the TPE-PDT efficacy of PORBSNs-mannose on human tumor. For this, zebrafish embryos were injected with MDA-MB-231 human cancer cells stained with (red) Dil staining. After 2 days, the colored tumors developed. We have demonstrated that the intravenous injection of PORBSNs-mannose allowed targeting the tumor as confirmed by the reduction of the tumor size after irradiation. The tumor sizes increase in all other conditions such as laser alone or injection alone, demonstrating the efficiency of the focused irradiation of tumor area with a pulsed laser when animals were treated with targeted nanoparticles.

Finally, in this study, we decided to go further in the description of the mechanism involved in tumor reduction. Although we have demonstrated the elevation of ROS production in cancer cells, we decided to show the involvement of caspases apoptotic pathway during the TPE-PDT in zebrafish embryos bearing tumor. Our experiments described that the tumor disappearance was concomitant and colocalized with an important increase in caspase-3 level. This is consistent with what is known about the involvement of apoptotic pathways in PDT-induced cell death.33,36

More generally, the use of BSNs for cancer targeting is still at the very beginning, and we are convinced that small-sized BSNs...
represents a breakthrough technology in the field of TPE-PDT for cancer treatment.

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**CONFLICT OF INTEREST**

The authors declare that they have no conflicts of interest.

**AUTHORS’ CONTRIBUTION**


**DATA AVAILABILITY STATEMENT**

Corresponding author promises to provide the data when requested.

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