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Self-Assemblies of Fe₃O₄ Nanocrystals: Toward Nanoscale Precision of Photothermal Effects in the Tumor Microenvironment

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 Fe_3O_4 nanocrystals were self-assembled into two different conformations: colloidosome and supraball, that confer them with distinct properties determining their photo-induced heating capacities. We asses these selfassemblies for photothermal therapy, an adjuvant strategy for tumor therapy. The tumor microenvironment is an heterogeneous ecosystem including immune cells and the extracellular matrix. The interactions between photothermal therapy agents and the different components of the tumor microenvironment determine the outcome of this therapy. In this study we decorticate the fate of both colloidosomes and supraballs within the tumor microenvironment in comparison to their Fe_3O_4 nanocrystal building blocks. We highlight how these two hybrids self-assemblies target different compartments of the tumor microenvironment and trigger local photothermal damages that are inaccessible for isolated nanocrystals and not predicted by global temperature measurements.

1. Introduction

Self-organization of entities in 2D and 3D structures is a process frequently found in nature. For example, sodium and chloride atoms self-organize in a highly ordered structure to form NaCl salt, and phospholipids self-assemble in a lipid bilayer to form the cell membrane. Inspired by this natural phenomenon, chemists have developed methods to build artificial colloidal crystals of nanocrystals called supra/super-crystals. ^[1-4] The chemical and physical properties of such crystalline structures are neither those of the building blocks nor those of the bulk materials. They are due to either dipolar interactions between the building blocks (collective properties) ^[5,6] or related to the crystalline structures (intrinsic properties). ^[7-9] These unique features of self-organized nanostructures open new avenues for medical purposes, but have been poorly explored so far. For particular applications such as photoexcitation and photothermal therapy (PTT), the fine-tuning of the optical properties as well as the control of the interactions between the nanomaterial and the biological environment is of upmost importance.

Recently, a new concept to lock iron oxide nanocrystals at the liquid-liquid interface has been developed: a "Janus bilayer" shell on the ferrite nanocrystals coated with aliphatic ligand can adsorb at the liquid-liquid interface inducing the formation of flexible colloidosomes. ^[10] This approach avoids complicated procedures, such as those found in nanocrystal polymersomes. ^[11,12] When nanocrystals are not locked at the liquid-liquid interface, solid fcc spherical supraballs, are produced. Fcc spherical supracrystals have been previously obtained

with either 6 nm cobalt iron oxide ^[13] or colloidal silica particles. ^[14] The collective properties of these new hybrid suprastructures (colloidosomes and supraballs) are still in their infancy and seem to depend on the medium of application. For example, due to the high penetration depth of visible light in fcc supracrystals dispersed in aqueous solution and effective accumulation of energy in the internal reservoir formed by the coating agents, these structures can serve as very efficient nanoheaters. ^[15] This opens a broad range of potential applications in energy release including solar energy and biomedicine.

Photothermal therapy (PTT) elicit new clinical trials as adjuvant or neo-adjuvant strategy to be applied in chemo-resistant and difficult to treat tumors that have to be reduced before surgery. Photoactivable nanomaterials including plasmonic metallic nanoparticles (i.e. gold nanoshells), ^[16] carbon absorbing nanomaterials (i.e. carbon nanotubes, graphene and analogues) or iron oxide nanoparticles ^[17] are remotely activated by a laser which penetrate the tissue generating a controlled hyperthermia in the target region. In order to be usable in clinics, a PTT nanoagent must absorb optical wavelengths in the tissue transparency windows once in the tumor environment, and fulfil criteria of low toxicity to healthy organs, easy functionalization and scale-up, and favorable biodistribution and life cycle in vivo. [18] However, there is plenty of room for improvement of their heating capacity and fine control of the biological outcomes of nano-hyperthermia that strongly depend on the interactions of nanostructures with the local biological environments. In this respect, iron oxide nanoparticles are particularly attractive due to their natural metabolization by endogenous proteins, high biocompatibility and clinical approval of several formulations that are combined with unique superparamagnetic properties for magnetic resonance imaging, photoacoustic tomography, magnetic manipulation or magnetic hyperthermia as well as iron related redox reactivity and light to heat conversion. Whereas iron oxide nanoparticles are susceptible to both magnetic and light activation to produce local heating, the photothermal strategy has proved more efficient to induce tumor hyperthermia at low dosage. [19-25]

In solid tumors, nanoparticles encountered not only tumor cells but different components of the tumor microenvironment that include stromal cells (among which we can find fibroblasts or immune cells) and a copious lattice of extracellular matrix. The intimate interactions of the nanomaterials with the heterogeneous ecosystem of the tumor will dictate their tumor penetration, local distribution of nanoparticles, local heating efficiency, and the ultimate biological damages and mechanisms of resistance that are induced. Nevertheless, little is known on how nanomaterials interact with the tumor microenvironment and how their structural properties can modulate these interactions, their heating capacity and the PTT-induced biological outcomes. Particularly, the use of self-assembled nanocrystals forming organized suprastructures is still a blank field despite their attractive properties. In a recent paper, we have shown that flexible colloidosomes, rigid supraballs and their isolated building blocks differently interact with tumor cells, resulting in distinct intracellular distribution. Remarkably, self-assemblies allow to optimize the lysosomal density and localization of iron oxide nanocrystals and to maintain nanocrystal organization within the lysosomes.^[26] Whether the organization of the self-assemblies translates into specific photothermal properties in biological environment is currently unknown.

The present study aims to evaluate how the specific features of self-organized nanostructures can influence their efficiency as photothermal therapy agents in different biological models recapitulating the tumor microenvironment, including cellular models, *ex vivo* and *in vivo* tumor models. We highlight the remarkable result that organic inorganic self-assemblies of nanocrystals can target different components of the tumor microenvironment with respect to individual nanocrystals. Moreover, self-assemblies generate local thermal damages that are inaccessible for isolated nanocrystals and not predicted by global temperature measurements.

2. Results and Discussion

2.1. Ferrite Colloidosomes and Supraballs

Hydrophobic ferrite nanocrystals (from now on Fe₃O₄ NCs) coated with oleic acid (OA, [CH₃-(CH₂)₇-CH=CH-(CH₂)₇-COOH]) are characterized by an inverted spinel structure. By ligand exchange process Fe₃O₄ NCs are coated with dopamine and dispersed in aqueous solution. Whatever the coating agent, the average diameter of Fe₃O₄ NCs are 10 ± 1 nm (Figure 1A and Figure S1A, Supporting Information).

By mixing Fe_3O_4 NCs coated with OA dispersed in chloroform and dodecyl trimethylammonium bromide (DTAB, $[C_{12}H_{25}N(CH_3)_3^+Br^-]$) dissolved in aqueous solution, two suprastructures were produced. The overall procedure remained the same except the presence or not of octadecene (ODE, $[CH_3^-(CH_2)_{15}^-CH^=CH_2]$) in chloroform.

In presence of ODE, hollow self-assembled Fe_3O_4 NCs structures, named colloidosomes, consist on one or a few layers of Fe_3O_4 NCs (less than 4) with 170 ± 90 nm as average diameter (Figure 1 and Figure S1B, Supporting Information). The inner region is made by the tail part of OA strongly bound, via the carboxyl group, to the surface of the Fe_3O_4 NCs. In contrast, the outer shell is composed of DTAB, ODE and OA. Strong hydro- phobic interactions between the alkyl chains as well as the double bonds of OA and ODE favored the dispersion of suprastructures in aqueous solution.

In absence of ODE during the mixture of the two solutions mentioned above, instead of colloidosomes, spherical solid self-assemblies of Fe₃O₄ NCs in fcc crystalline structures called supraballs with an average diameter of 69 \pm 15 nm were produced (Figure 1C and S1C). The contrast change in the TEM image (Figure 1C) is characteristic of the step edge structure of the assembled Fe₃O₄ NCs, as previously observed for fcc cobalt iron oxide and silicaparticles. ^[13, 14]

The nanomechanical properties of these suprastructures, load-depended in aqueous solution, show that colloidosomes have a higher flexibility, deformation, stiffness and elastic moduli than supraballs. Presence of

ODE molecules in the shell of the colloidosome induces an increase in deformation and adhesion forces at the edges.^[26]

Below we compare the photothermal properties of Fe_3O_4 NCs and either colloidosomes or supraballs, all dispersed in aqueous solution. Note these nanostructures keep their integrities in size, size distribution, structure and pH in the range from 5 to 8 for more than three years.

2.2. Suprastructures Photothermal Properties: Apparent Contra- diction between the Global Heating and Cell Death

To evaluate the photo-induced heating properties of Fe_3O_4 NCs, colloidosomes and supraballs, we first compared the absorption spectra of the suprastructures and dispersed Fe_3O_4 NCs in aqueous solution. Unexpectedly, the suprastructure absorption spectra considerably dropped compared to their building blocks (Figure S2, Supporting Information). The photo-activated heating of the hybrid structures or dispersed Fe_3O_4 NCs was obtained by subjecting 100 µL of suspension at fixed iron con- centration (150 µg mL⁻¹) to an 808 nm laser power (1 W cm⁻²) for 5 min. The temperature increment was the largest for supraballs (10.9 °C), followed by Fe_3O_4 NCs (8.8 °C), whereas the colloidosomes heated by less than 2 °C (Figure S3, Supporting Information). According to the drop of the suprastructure absorption spectra compared to that of dispersed Fe_3O_4 NCs, we could expect a lower temperature increment for both hybrids compared to Fe_3O_4 NCs. However, it was previously observed that Fe_3O_4 NCs self-assembled in fcc superlattices act as nanoheaters embedded in a thermal reservoir formed by the coating agents.^[15] This could explain the difference in the temperature increment between supraballs (spherical fcc supracrystals) and colloidosomes (closed structure of one or a few layers of Fe_3O_4 NCs). The rather large value obtained with Fe_3O_4 NCs compared to colloidosomes can be explained by the large difference in the number of absorbed photons at 808 nm.



Figure 1. CryoTEM images of A) Fe_3O_4 NCs, B) colloidosomes, and C) supraballs.

In our previous paper, ^[26] by using self-assembled nanostructures such as colloidosomes and supraballs, we developed a new concept to investigate their intracellular fate. To evaluate the effect of internalization by tumor cells on the global heating efficiency of Fe₃O₄ NCs, colloidosomes and supraballs, A431 human epidermoid carcinoma cells were incubated with the nanostructures at an iron concentration of 28 μ g/mL for 24 h (Figure 2A).^[28] The cells were then washed carefully to discard the excess of Fe₃O₄ NCs that had not been internalized and one million of cells were pelleted and resuspended in 100 μ L of culture medium before irradiation with an 808 nm laser in the same experimental conditions as for the nanostructures in suspension. Dispersed Fe₃O₄ NCs heating efficiency was significantly higher to that of supraballs and colloidosomes once internalized in cells. From initial temperature of 25°C, ^[29],cells reach a plateau temperature (in about 200 s) of 38.4, 33.5 and 28 °C for Fe₃O₄ NCs, supraballs and colloidosomes, respectively (Figure 2B-D). Colloidosomes are therefore the least efficient for a global heating when suspended in water or internalized in cells.



Figure 2. Evaluation of $\text{Fe}_{3}\text{O}_{4}$ NCs, colloidosomes and supraballs heating capacity in A431 cells. A) Scheme of the experimental setup to evaluate the global effect of laser irradiation on cell pellets. After 10 min laser irradiation of the nanoparticle-loaded cell pellet, cells are redeposited in cell culture flasks for 4 h before evaluating the cell death. B) Temperature increase in a pellet of 1 million cells having internalized $\text{Fe}_{3}\text{O}_{4}$ NCs, colloidosomes and supraballs and dispersed in 100 µL of PBS upon irradiation with an 808 nm laser (1 W cm⁻²). C) Infrared thermal images of cell pellet captured at 5 min of 808 nm laser irradiation. D) Temperature increase of cell pellets compared to that of aqueous suspension of $\text{Fe}_{3}\text{O}_{4}$ NCs, colloidosomes and supraballs at two different concentrations [Fe] = 150 and 1000 µg mL⁻¹. The concentration of Fe in cell pellet is calculated from ICP-AES quantifications. E) Percentage of cell necrosis and apoptosis upon laser exposition (or not) for 10 min of cell pellet described above. Results shown as mean ± SEM (20–30 images per condition, n = 3). One-way ANOVA, Kruskal–Walis, ****p*-value < 0.001. F) Representative fluorescence images of A431 cells 4 h after laser irradiation. Cells were stained with Hoechst 33342 (cell nuclei, blue), YO-PRO (apoptotic cells) and propidium iodide (necrotic cells). Scale bar = 50 µm.



Figure 3. Evaluation of the local cellular damages induced by Fe_3O_4 NCs, colloidosomes and supraball-mediated PTT in cell monolayers 4 h post- irradiation. A) Scheme of the experimental setup to evaluate the local effects. A431 cells were incubated overnight with Fe_3O_4 NCs, colloidosomes and supraballs, and after rinsing, they were irradiated as monolayer with an 808 nm laser for 10 min. The cell damaged was evaluated 4 h post-irradiation. B) Percentage of cell necrosis and apoptosis caused by Fe_3O_4 NCs, colloidosomes and supraballs-mediated PTT in cell monolayers. Results shown as mean \pm SEM (20–30 images per condition, n = 6). One-way ANOVA, Kruskal–Walis, **p*-value < 0.05, ***p*-value < 0.01, ****p*-value < 0.001. C) Representative fluorescence images of A431 cells. Cells were stained with Hoechst 33342 (cell nuclei, blue), YO-PRO (apoptotic cells) and propidium iodide (necrotic cells). Scale bar = 50 µm.

Apart from this global heating efficiency of nanostructures submitted to irradiation, the important outcome for therapy is the biological effects of Fe₃O₄ NCs-mediated heating. We thus quantified the cellular damages related to PTT, that might arise from the global heating of the tissue or cell pellet, or from more local thermal damages induced in the vicinity of the heat nanosources. Indeed, it has been shown that the temperature increase close to the heat nanosources is much higher than the macroscopic temperature in the surrounding medium. ^[30,31] For this purpose, we evaluate thermal damages on cells in two different laser irradiation set-ups: i) in the pellet of NC-loaded cells (as described above) which combines local intracellular heating with global temperature increase in the pellet measured by thermal imaging, and ii) on monolayers of cells having internalized the Fe₃O₄ NCs for which the global temperature is thermalized by the particle-free culture medium at 37°C (Figure 3A). In the latter set-up, only local effects of intracellular heating can be expected.

Cell apoptosis and necrosis were assessed through the evaluation of the membrane permeability to fluorescent dyes. Hoechst 33342 stains the condensed chromatin of live cells, this allows the quantification of all cells. YO-PROTM dye can enter cells under apoptosis whilst, propidium iodide cannot, it only stains dead necrotic cells. The staining pattern resulting from the simultaneous use of these three dyes allows to distinguish alive, apoptotic and necrotic cells populations by fluorescence microscopy represented in Figures 2F and 3C and quantified in Figures 2E&3B. As previous studies demonstrated the detection of cell death apoptosis 4-5 hours post

irradiation, ^[32] cells that have been pelleted and irradiated were then put back in culture for 4 hours before cell death assessment (Figure 2A). Cells having internalized the different nanostructures without laser irradiation did not show any significant enhancement in apoptosis or necrosis level in comparison to the control cells, irradiated or not, showing that the nanostructures or the laser exposure did not affect cell viability itself. This confirms our previous study ^[26] investigating the fate of the same nanostructures up to 8 days post-labeling, showing no significant toxicity in A431 cells up to concentration of 112 μ g/mL in absence of irradiation.

Cells having internalized isolated Fe₃O₄ NCs show slightly enhanced level of apoptosis and necrosis after irradiation, but the difference with non-irradiated cells is not statistically significant. In contrast, colloidosome and supraball-loaded cells feature a significant enhancement of the apoptosis (more than 15% of total cell number) and necrosis level (~15%) after irradiation. These results are surprising considering the lower global temperature of the cell pellet for supraballs and colloidosomes in comparison to Fe₃O₄ NCs (Figures 2B-C). This suggests that supraballs and to a larger extent colloidosomes could induce local thermal damages independently of the global temperature increment in the cell pellet. This has to be related to previous data ^[15] from which it was demonstrated that the light absorption in such nanostructures, dispersed in aqueous solution, initiates a chain energy transfer having various degree of freedom coupled together. In vitro, the efficiency of supraballs is expected to be larger than colloidosomes. However, in cells, the colloidosome shape is not retained and the nanocrystals self -assembled in large domains and consequently in first approximation we could assume, but not quantify, that energy penetration is almost an even larger for colloidosomes than supraballs.

To isolate thermal damages that arise from intracellular local heating, thermalized cell monolayers were exposed to the laser and cell death was quantified in the same conditions as for the cell pellet (Figure 3A). There was no effect of apoptosis or necrosis in absence of laser irradiation, but we observed significant increase of apoptosis/ necrosis level after irradiation for the three types of nanostructures. The level of cell damage induced by direct irradiation of cell monolayers was significantly higher as compared to that in pellet form, although no macroscopic heating can be measured. Even though the laser irradiation experienced by each cell can be higher than in the cell pellet as there is no light screening by cells and nanoparticles absorption, we expect in this set up only local effect of intracellular nanoheaters. Remarkably, despite colloidosomes have low global heating performance in aqueous solution and even lower in cells, colloidosome-mediated intracellular local absorption induces the highest levels of cell damage compared to Fe_3O_4 NCs and to supraballs (Figure 3B, C). Fe_3O_4 NCs, despite exhibiting higher heating efficiency both in aqueous solution and in cell pellet, induce less cellular damage upon irradiation of cell monolayers. Overall, the two self-assemblies of Fe₃O₄ NCs clearly induce massive cell death: 74% and 57% of cell damage (both necrosis and apoptosis) induced by colloidosome and supraball-mediated PTT, respectively (Figure 3B), in com- parison, to less than 35% for dispersed Fe₃O₄ NCs. This experi- ment clearly demonstrates that self-assemblies of Fe₃O₄ NCs, once internalized by cells, can elicit massive local damages that provoke cell apoptosis and necrosis without global heating of the environment.

At this point we asked the following question: What is the origin of the apparent contradiction between the global heating efficiency and the biological outcome on cells? To answer such question, we highlighted the intracellular distribution of the Fe₃O₄ NCs. In our previous study, ^[26] we described important differences in the intracellular fate of Fe₃O₄ NCs internalized as dispersed Fe₃O₄ NCs or as self-assemblies in the form of colloidosomes and supraballs up to 8 days post labeling (see Figure 4 for intracellular distribution at 24 h). Upon cell internalization, dispersed Fe₃O₄ NCs tend to aggregate into endosomes that fuse with lysosomes, forming uncontrolled fractal structures similar to those obtained by diffusion-limited aggregation. In contrast, supraballs and colloidosomes conserve their organized structures over at least 8 days once internalized by tumor cells demonstrating that self-organization of Fe₃O₄ NCs could withstand intracellular trafficking and the harsh acidic environment of lysosomes. In addition, supraballs and colloidosomes self-assemblies maximize the intralysosomal density of Fe₃O₄ NCs (surface density of 52% and 34%, respectively) in comparison to 4% for isolated Fe₃O₄ NCs. Moreover, looking at the spatial distribution of Fe₃O₄ NCs within the lysosomes, a much higher percentage of Fe₃O₄ NCs were located within 100 nm of the lysosomal membrane after colloidosome internalization (up to 48%) compared to that of dispersed Fe₃O₄ NCs (9%) and to a minor extend compared to supraballs (20%), as illustrated in Figure 4. ^[26] Such local accumulation of Fe₃O₄ NCs close to the lysosome membrane could explain that larger local damages could be obtained with self-assemblies in comparison to dispersed Fe₃O₄ NCs. In addition to the spatial density of Fe₃O₄ NCs, their ordered structure into the lysosome, inherited from the self-organization of the assembly, might favor an accumulation of energy within these structures. Indeed, as demonstrated earlier by ultra-fast optical spectroscopy and semiclassical modeling, ^[15] the dilution of the metallic phase in the hybrid self-assemblies enable to increase the penetration depth of visible light in comparison to homogeneous metallic NCs of similar size, thus enabling an energy load in the assembly. The same study revealed that coating agents of Fe₃O₄ NCs act at short time as a reservoir of energy due to the tiny distance between nanocrystals and thermal transfer that propagates preferentially within the assembly. This reservoir effect and thermal resistance of the hydrophobic organic phase consequently induce an increase of the localized heating in the self-assemblies compared to the fractal aggregates of Fe₃O₄ NCs. Hence, we hypothesize that the optimal accumulation and local density of Fe₃O₄ NCs in lysosomes, their sustained compacity in lysosome and the affinity of the colloidosome shell for the lysosomal membrane can collectively explain the enhanced local damages that were observed upon irradiation, inducing massive cell death. The role of lipid bilayers and organic coating as a thermal barrier that favor local accumulation of energy has been demonstrated before using nano-thermometry.^[33] Here, we observe a direct biological consequence of the energy accumulation in the lysosome membrane due to the close proximity of the lysosome membrane to the absorbing Fe_3O_4 NCs in colloidosomes and supraballs. This accumulation may induce lysosome swelling and membrane permeabilisation much more efficiently than for isolated Fe₃O₄ NCs, without significant global heating. We thus conclude that the optimization of Fe₃O₄ NC density in lysosomes and proximity to the lysosome membrane, together with the high energy accumulation within the hydrophobic shell of colloidosomes or the hydrophobic core of the supraballs enhance local hot spots inducing cell death. Our study clearly demonstrates that cell damages are not directly related to the global heating of the environment but rather to the local energy accumulation that depends on ultrastructural distribution of Fe₃O₄ NCs among themselves and within the lysosome. These data agree with previous study showing that cancer cells can be particularly sensitive to agents which induce lysosomal cell death through lysosome membrane permeabilisation or disruption. ^[34] It results in the activation of apoptosis signaling pathways and contributes to the amplification of death signals.^[35] Lysosome membrane permeabilization can be triggered by a wide variety of agents, one of them being local heating.^[36,37] In addition, we have quantified in our previous paper ^[26] that colloidosomes induce a significant enhancement of the lysosome size at 24 h post labeling without inducing toxicity in absence of irradiation. Thus following laser stimulation, the local heating might trigger enhanced lysosome swelling, affecting more easily the lysosome integrity than with supraballs and with isolated nanocrystals, as observed in Figure 4B.



Figure 4. Colloidosome and supraball intracellular distribution in A431 cells. TEM images of A) Fe_3O_4 NC, B) colloidosome and C) supraball-loaded cells 24 h after incubation illustrating the distinct intralysosomal distribution of nanocrystals.

2.3. Suprastructures: Photothermal Properties in the In Vivo Tumor Microenvironment

To test the feasibility of photothermal therapy mediated by self-assembled nanostructures in solid tumor, Fe₃O₄ NCs, colloidosomes and supraballs were injected intratumorally (i.t.) in subcutaneous tumors of murine breast cancer model MET-1. ^[38] The i.t. administration was chosen for two reasons. Firstly, it is a clinically relevant option for tumors that should be shrink before surgery to improve local control (neoadjuvant treatment). Percutaneous or endoscopic administration of nanoparticles has been achieved in many solid tumors such as soft tissue sarcoma, breast, pancreatic or gastrointestinal cancers and endoscopic and intraluminal light illumination

is also feasible through the same procedures.^[39-42] Secondly the local administration of nanoparticles avoids the massive off-target accumulation of nanoparticles associated with intravenous or intra-arterial injection, minimize systemic toxicity and maximize the tumor concentration of nano-heaters. It is also the best option for comparing nanostructures that may have very different biodistribution and access to the tumor when injected intravenously. Here, the same mass of iron ([Fe]= $150 \ \mu g$ in 50 μL injected volume) was infused in tumors of approximately 150-200 mm³ which allowed a head to head comparison of the different nanostructure distribution in the tumor microenvironment and the outcome of light exposition. Injection of nanomaterials was performed in a unique injection point with a very slow infusion through 27 G needle to maximize the distribution of nanostructures which were let for 24 h to diffuse and to be processed by cells of the tumor microenvironment before irradiation. 24 h after i.t. administration, tumors were exposed for 15 minutes to an 808 nm laser at a power of 2 W/cm². The tumors' surface temperature was monitored with a near infrared camera (Figure 5A). A significant increase in the tumors' surface temperature was observed with all nanostructures (plateau temperature of T=45, 44 and 43°C for supraballs, Fe₃O₄ NCs and colloidosomes) (Figure 5B) in comparison to the laser-exposed non-injected tumors. 24 hours after laser irradiation, tumors were harvested and the local effect of Fe₃O₄ NCs, colloidosomes and supraballs-mediated PTT was evaluated using histology. Tumors were stained with Perls' Prussian Blue staining that allowed the detection of iron in tissue samples, in order to localize the Fe_3O_4 NCs within the tissue. They were also stained with hematoxylin eosin saffron (HES) staining, hematoxylin stained the cell nuclei in blue, eosin stains the extracellular matrix and the cytoplasm in pink, whilst saffron allows the highlighting of collagen fibers (stained in yellow/orange). This trichromatic stain allows a general evaluation of a tissue sample and identification of anomalies such as the presence of necrotic regions. The presence of necrotic regions was identified for each of the conditions (Figures 5C-D). The basal level of tumor necrosis without laser exposure was about 13% regardless of the nanostructures injected or not. However, we observed striking differences in the PTT-induced necrosis level depending on the nanostructures inside the tumors. Even more surprisingly, the necrosis level did not correlate with the tumor temperature increase. Actually, the largest necrosis extension (64%) was observed for colloidosome-treated tumors which however heat at only 43°C for less than 15 min under laser exposure. Supraball-mediated PTT at the same laser power induced a 50% necrosis extension for a surface temperature of 45°C (Figures 5C-5D). Fe₃O₄ NCs on the other side, induced a much lower level of tissue damage (24%) while the temperature plateaued to 44°C. This evidenced again that the level of global hyperthermia applied to a given tumor does not necessary translate to a higher level of cellular damage of the tumor tissue. Importantly we observe that hybrids (colloidosomes and supraballs) induce a significant increase in tissue necrosis extension (>50% of the tissue) after a single laser exposition and 15 min of mild hyperthermia, while the biological outcome of their individual building blocks in similar experimental conditions and macroscopic temperature increase is not significant. These results at the tissue level in tumors fully correlates with the results obtained in the experiments described above at the cellular level, suggesting superior toxicity of intracellular heating mediated by self-assemblies in the tumor tissue.



Figure 5. Colloidosome and supraball-mediated photothermal therapy in vivo. A) In vivo infrared thermal images of tumors injected with 150 µg colloidosomes or supraballs and exposed 15 min to an 808 nm laser at 2 W cm (n = 5). C) Quantification of the percentage of necrotic areas in MET-1 tumor in each of the tested conditions (n = 5). One-way ANOVA, Kruskal–Walis, *p-value < 0.05, **p-value < 0.01. D) Histological (Perls and HES staining) analysis of Fe₃O₄NCs, colloidosome and supraball injected MET-1 tumors 24 h after laser irradiation or without laser exposure. Scale bar = 50 µm. Nanoparticles appearing in blue with Perls staining are located in the tumor. Necrotic areas underlined by dashed line can be observed in tumors treated with laser close to NC localizations.

2.4. Suprastructures Modulate the Distribution of Fe₃O₄ NCs in the Tumor Microenvironment

Tumors do not consist exclusively on tumor cells, but on a complex ecosystem including stromal cells (endothelial cells, fibroblasts and immune cells) and extracellular matrix (ECM) composed of a multitude of biomolecules including collagen and hyaluronic acid, in what is called the tumor microenvironment.^[43] The role of the heterogeneity of the tumor microenvironment in chemoresistance has been evidenced in the last decade ^[44]

but somewhat overlooked in the field of nanoparticle-mediated therapies since the precise nanoparticle localization and local effects have been barely investigated.^[45] Considering that the components of the tumor might interacts differently with the nanostructures, we thought to investigate the differences in tissue distribution and local photothermal effects that could arise specifically from the hybrid self-assemblies. Importantly, we could observe striking differences one day after i.t. injection in the MET-1 breast tumor environment (Figure 6). As already mentioned, Perls staining detect iron in blue, the hematoxylin stains cell nuclei blue, eosin stains the extracellular matrix and cytoplasm in pink, Fe₃O₄ NCs appears in yellow/orange on HES staining and F4/80 detect macrophages in brown. When isolated Fe_3O_4 NCs were injected in tumors (Figure 6, left), they mainly localized in the extracellular space, in close contact with the ECM fibers. Such localization along collagen fibers was previously observed for iron oxide nanocubes of 20 nm.^[46] In addition, a recruitment of F4/80 positive macrophages (indicated by blue arrows in Figure 6) is induced in the vicinities of Fe₃O₄ NCs. Contrastingly colloidosomes are mainly internalized by tumor cells (Figure 6, middle), whilst supraball location is both intracellular and extracellular (Figure 6, right). This revealed that the different nanostructure architectures severely impact the nanostructure distribution in the heterogeneous tumor environment. Particularly the hybrids, and more particularly colloidosomes have a more potent tropism for tumor cells in comparison to their dispersed building blocks (Fe₃O₄ NCs) which distributed mostly in between collagen fibers in this breast tumor model. This is an important finding to explain the extended necrosis damage inflicted by colloidosome and supraballmediated PTT in comparison to the insignificant effect of dispersed Fe₃O₄ NCs, despite similar temperature increase. Colloidosomes and to a lesser extent supraballs could effectively induce deleterious local heating within tumor cells due to their specific accumulation, while Fe₃O₄ NCs, in addition to a lower intracellular toxicity after irradiation, were mostly located extracellularly with lesser impact on tumor cells.

To decorticate further the interactions of the nanostructures with the heterogeneous populations of the tumor microenvironment, we compared *in vitro* the internalization of dispersed Fe₃O₄ NCs, colloidosomes and supraballs with separated monoculture of tumor cells (A431 – human epidermoid carcinoma cells), macrophages (RAW 264.7- murine macrophages) and fibroblasts (hTERT-HSC-human immortalized liver fibroblasts). Iron quantification by ICP-AES displayed in Figure 7 shows that supraballs could transfer more iron in macrophages than colloidosomes and dispersed Fe₃O₄ NCs. In contrast, iron capture in tumor cells was higher with colloidosomes than with supraballs and dispersed Fe₃O₄ NCs. Fibroblasts also present higher uptake of iron with the colloidosomes and to a lesser extent with supraballs in comparison with dispersed Fe₃O₄ NCs that show more than ten times smaller uptake.

Our comparative study emphasizes the poor efficacy of dispersed Fe_3O_4 NCs to accumulate into tumor cells or into fibroblasts, even if these cells are separately exposed to the Fe_3O_4 NCs in absence of ECM. Importantly, the only cell type that accumulates significant amount of dispersed Fe_3O_4 NCs (>50 pg) are the macrophages in line with the histological observations in tumors. Remarkably the self-assembly of Fe_3O_4 NCs within prescribed suprastructures can reverse this trend since colloidosomes are also substantially up taken by tumor cells and fibroblasts. Such differential cell internalization depending both on cell types and nanostructure architectures can be interpreted in view of their different mechanical properties, such as stiffness and deformability. Macrophages are key actors of immunity and first line of defense against pathogens, xenobiotics or senescent cells. It has been reported that macrophages can sense the rigidity of phagocytic targets, such as red blood cells that become more rigid after senescence or in certain disease conditions (thalassemia, sickle cells). ^[47] A number of studies have shown that stiffer particles are internalized to a higher extent than softer particles by macrophages and other immune cells. ^[48,49] Our results show that stiffer supraballs are also more internalized by macrophages than colloidosomes with slightly lower Young's modulus and a more deformable structure. ^[27]



Figure 6. Histological (Perls and HES staining) and immunological (F4/80 labeling) analysis of MET-1 tumors 24 h after intratumoral injection of $\text{Fe}_{3}\text{O}_{4}$ NCs, colloidosomes and supraballs. Insets display magnification of the gray square. Iron appears in blue with Perls staining and in yellow/ orange in HES and F4/80 images. White arrows indicate ECM fibers along which most $\text{Fe}_{3}\text{O}_{4}$ NCs are located. Gray arrows indicate the recruitment of macrophages (F4/80 brown labeling). Colloidosomes have mostly intracellular localization (see the inset on Perls staining) and supraballs have both intracellular and extracellular localizations.

One explanation for increased uptake of supraballs by phagocytic cells is that colloidosomes may undergo deformation in response to phagocytosis forces exerted by the cells which locally decreases their radius of curvature and makes phagocytosis more energetically costly and more difficult to achieve than with rigid undeformable supraballs. ^[50] In contrast to macrophages, the comparison of tumor cell uptake of soft/hard particles has shown complex findings without general trend emerging. ^[51,52] In this case, epidermoid carcinoma cells show a preference for soft and deformable colloidosomes. This is also in line with our finding in the

in vivo MET-1 breast tumor model confirming higher uptake of tumor cells for colloidosomes. To sum up, both in the heterogeneous microenvironment of tumors and in conditions in which only one cellular component is exposed to the nanostructures, the self-assembled colloidosomes have the unique potential to maximize their accumulation by tumor cells and fibroblasts in comparison to ECM and macrophages. This is the first demonstration that the self-organization of Fe₃O₄ NCs and the resulting mechanical properties can be used to modulate the distribution of Fe₃O₄ NCs in the tumor microenvironment.



Figure 7. Iron quantification measured by ICP-AES in tumor cells (A431 cell line), macrophages (RAW 264.7 cell line) and fibroblasts (hTERT-HSC immortalized line) after 24 h exposition to Fe_3O_4 NCs colloidosomes or supraballs at [Fe] = 28 µg mL⁻¹ (*n* = 3). One-way ANOVA, Kruskal–Walis, * *p*-value < 0.05, ***p*-value < 0.01, *** *p*-value < 0.001.

The ECM is another important component of the tumor stroma that is differently expressed and organized depending on the origin and progression of the tumor. In order to investigate the interaction between the nanostructures and the ECM, we set up an in vitro model consisting of fibroblast sheets secreting ECM over stimulation with ascorbic acid. ^[53] The abundant ECM fibers surrounding fibroblasts can be observed on TEM image in Figure 8A. These fibroblast sheets were exposed to Fe₃O₄ NCs, colloidosomes or supraballs overnight, followed by careful washing. As in the in vivo tumor, Fe₃O₄ NCs were mainly located along the ECM in between fibroblasts (Figure 8B). The colloidosomes were all internalized by fibroblasts and were absent from ECM (Figure 8C). Supraballs could be found in both compartments (Figure 8D). These results confirm that in the hybrid ECM/cellular environment, Fe₃O₄ NCs are barely internalized by fibroblasts, whilst colloidosomes show the opposite behavior. Notably, supraballs are less internalized by fibroblasts than colloidosomes. The mechanical properties of colloidosomes featuring lower Young modulus/stiffness and larger flexibility than supraballs may favor internalization by fibroblasts, while supraballs are more easily trapped in ECM fibers, as well as isolated Fe₃O₄ NCs. Such observations on fibroblast sheets nicely correlate with those shown previously in the complex microenvironment of the tumor (Figure 6) in which Fe₃O₄ NCs were mainly located in the ECM, colloidosomes internalized in the tumor cells and supraballs both located in intracellular and extracellular compartments.



Figure 8. TEM images on 70 nm microtome sections of Fe_3O_4 NCs, colloidosome or supraballs distribution in the fibroblast/ECM in vitro model. The fibroblasts are highlighted in yellow in the lower magnification images. Images on the right are magnification of the white quadrant on image on the left. A) Control fibroblast sheet with dense ECM. B) Fe_3O_4 NCs distribution on fibroblast-ECM in vitro model, Fe_3O_4 NCs mainly accumulate in the ECM compartment. C) Colloidosomes distribution on fibroblast-ECM in vitro model, mainly internalized in fibroblasts. D) Supraball distribution on fibroblast-ECM in vitro model, distributed both within fibroblasts and within the ECM.

2.5. Suprastructures: Photothermal Effects on the Tumor Extracellular Matrix

Due to the important role of ECM in promoting tumor progression, chemoresistance and immune exclusion, the ECM is recognized as a multicomponent target for advanced therapeutic strategies. Previous studies have showed that carbon-nanotube mediated PTT could induce the local denaturation of collagen fibers in tendon and in tumors.^[54] We thus tested the photothermal effects of the two hybrids and Fe₃O₄ NCs on ECM in different organotypic models. In order to visualize the direct effect of PTT on the ECM of real tumors, we used *ex vivo* fresh 350 μ m tumor slices incubated for 2 hours with Fe₃O₄ NCs, colloidosomes and supraballs (Figure 9B) and then exposed to 808 nm laser at 2 W/cm². The EGI-1 tumor model, a human bile duct desmoplastic cancer, was chosen for its abundant ECM (unlike the MET-1 tumor model previously used) and linear organization of collagen fibers that can be observed and quantified using second harmonic generation (SHG) two photon microscopy (Figure 9A). In comparison to the non-irradiated tumors, the amount of collagen SHG signal was

significantly reduced after irradiation of tumor slices exposed to colloidosomes (Figure 9C), but not to Fe₃O₄ NCs or supraballs. Analyses of SHG images with the CT-FIRE software ^[54] also show a significant decrease in collagen fiber length of tumors exposed to colloidosomes and irradiated (Figure 9D), that can be related to fiber denaturation induced by localized heating (Figure 9A, blue arrows).



Figure 9. Colloidosome-mediated PTT also induces collagen denaturation in EGI-1 tumor slices. A) Two photon microscopy SHG images of collagen network of EGI-1 (human bile duct) tumor slices incubated with Fe_3O_4 NCs, supraballs and colloidosomes and irradiated with an 808 nm laser. A clear reduction of collagen SHG signal can be observed after colloidosome-mediated PTT as well as a decrease of fiber length (blue arrows). Scale bar = 50 µm. B) Schema of the experimental set-up. C) Percentage of SHG collagen signal in the different conditions. D) Relative variation in collagen fiber length ($\Delta L/L$). Results are represented as mean ± SEM (One-way ANOVA, Kruskal–Wallis, *** p < 0.001, ** p < 0.01).

This de-structuration of collagen fibers on tumor slices by irradiated colloidosomes was confirmed in a tissue consisting of highly organized type I collagen, a fresh sheep tendon. Using the same protocol as for the tumor slices, colloidosomes induce the apparition of SHG signal voids in the dense collagen matrix of the tendon. This means that the collagen fibers have suffered a significant conformation change, leading to the loss of SHG properties, likely due to the local heat-induced denaturation of the collagen molecule. A significant increase in the presence and size of these areas devoid of SHG signals is observed when laser power is increased from 1 to 2 W/cm² (Figure 10A and 10B). However, collagen fibers remained intact after laser irradiation in the presence of Fe₃O₄ NCs and supraballs. To look at the collagen structure with a higher resolution, we performed Scanning Electron Microscopy (SEM) on the tendon before and after PTT (10 min at 2 W/cm²). The flattened structure of colloidosomes and the more compact and rigid supraballs are easily recognizable in SEM images of non-

irradiated tendon. After laser exposure, SEM observations confirm that colloidosome-mediated PTT induces the formation of "holes" on the dense collagen network (Figure 10C) close to colloidosomes. The same kind of holes are also seen in samples that had undergone supraball-mediated PTT (Figure 10D). This means that supraball-mediated PTT also induces collagen denaturation but to a lesser extent, probably not enough to be detected with SHG microscopy in these conditions. These experiments focusing on collagen matrix on tumor and tendon slices clearly reveal the colloidosomes as the most efficient light-activated nanoheaters to induce local denaturation of the collagen fibers in our experimental conditions.



Figure 10. Colloidosome-mediated PTT induces collagen denaturation. A) Two photon second harmonic generation (SHG) microscopy images of sheep tendon collagen alone (control) or incubated with Fe_3O_4 NCs, colloidosomes and supraballs and subsequently irradiated

with 808 nm laser a two different laser power (1 and 2 W cm⁻²). Collagen denaturation is observed as areas devoid of collagen SHG signals

within the dense collagen network (orange arrows). Scale bar = 50 μ m. B) Quantification of the presence and size of the collagen denatured sites for the different conditions. (One-way ANOVA, Kruskal–Wallis, ** *p* < 0.01). C) SEM images of fresh sheep tendon incubated with colloidosomes and irradiated with a NIR laser. Location of the colloidosomes is indicated with the white arrows. D) SEM images of supraballs deposited on the tendon and irradiated with a NIR laser. Location of the supraballs is indicated with the white arrows.

3. Conclusion

In this study we have focused on the photothermal and biological effects of Fe₃O₄ NC self-assembled into two different conformations: colloidosomes and supraballs. The different conformations confer them with unique properties, that govern both their local light-to-heat conversion as well as their tumoral tissue distribution in comparison to isolated nanocrystals. Remarkably, colloidosomes and supraballs are capable of inducing extended tumor necrosis after a single 15 minutes NIR in vivo laser exposition and mild hyperthermia in a breast cancer model in mice. This impressive tissue damage correlates with enhanced cell death on tumor cell suspension and monolayers and collagen destructuration in tumor slices of a desmoplastic tumor and in a sheep tendon. However, these biological outcomes of PTT mediated by self-assemblies do not correlate with the global temperature increase that is measured by infrared camera. Indeed, dispersed Fe₃O₄ NC exhibits higher global heating capacity than colloidosomes while their PTT biological effects are unsignificant. In contrast, colloidosomes and to a lesser extent supraballs induced potent and localized thermal damages both in tumor cells and collagen matrix. This is a unique demonstration that global heating measurements is neither representative of local energy deposition nor predictive of the biological outcomes of nanostructure-mediated PTT. Our findings emphasize that nanoparticle distribution and light to heat energy conversion must be controlled at the subtissular and subcellular scale in order to optimize the biological outcomes of "nanohyperthermia". This represents a paradigm shift in the field of nanoparticle-mediated hyperthermia.

Our study also reveal different phenomena that explain the outstanding properties of self-assemblies, in comparison to isolated nanocrystals, to optimize the NIR-induced local thermal damages in the biological environment: i) an enhanced energy accumulation in the hybrid self-assemblies with the organic phase playing the role of localized thermal resistance ^[15], ii) an optimized intralysosomal concentration of nanoheaters, iii) an optimized transfer of energy to the lysosome membrane that afford the affinity of colloidosomes to the cell membrane. All these features are direct consequences of the organization of the hybrid organic-inorganic self-assemblies and open new avenues for more precise control of photothermal conversion in the cellular and extracellular biological environment.

On top of these unique properties of hybrids as nanoradiators, we highlight very distinct distribution of the two hybrids and their dispersed building blocks (Fe₃O₄ NCs) in the complex tumor microenvironment, taken as a whole (intratumoral administration) or by separating its different components (tumor cells, stromal cells, extracellular matrix). Indeed, Fe₃O₄ NCs were mostly located in the ECM and macrophages, while supraballs were distributed partly in ECM and tumor cells. Owing to their different structure, flexibility and affinity for cell membrane, colloidosomes were shown to be more efficiently internalized by tumor cells than stromal cells and

to target the tumor cells even in the competitive multicomponent environment of the breast tumor model. This very efficient targeting of tumor cells by colloidosomes also correlate with enhanced PTT-induced necrosis in the tumor, while dispersed Fe₃O₄ NC have no significant effect. Here, we evidence that peculiar self-assemblies of nanocrystals allow for a better control of the tissular and intracellular distribution of the inorganic nanocrystals triggering different and more precise biological responses to the NIR irradiation. The complexity of the tumor microenvironment has been somehow overlooked in the research of nanomaterials for tumor therapy, which partly explain the low number of inorganic nanomedicines that have been translated into the clinics. Remarkably, using Fe₃O₄ NC or their self-assemblies as colloidosomes and supraballs allows us targeting different components of the heterogeneous tumor ecosystem which is an important asset to envision new strategies of personalized therapy. One possible way of exploiting these differences for more personalized and precision medicine would be choosing the correct nano-assembly as function of the therapy target and the peculiarity of the tumor microenvironment of the patient. This offers the opportunity to adapt the choice of nanostructures to the particularity of the actual tumor (e.g. fibrosis level, macrophage infiltration, importance of fibroblasts) and to the intended biological outcome (tumor cell death, ECM destructuration, fibroblast depletion, macrophage depletion ...) for seeking a better control of therapy.

4. Experimental Section

Cell culture. A431 human epidermoid carcinoma cells (ATCC, CRL-1555) were cultured in Roswell Park Memorial Institute medium (RPMI) supplemented with 10% fetal bovine serum and 1% penicillin and maintained at 37°C and 5% CO₂. Cells were seeding density and cell culture differed according the experiment performed. Murine macrophage-like cell line RAW264.7 (ATCC, TIB-71), hTERT-HSC derived from human activated immortalized hepatic stellate cells kindly provided by Dr. L. Aoudjehane (ICAN, Paris, France) and MET-1 cell line (obtained from Robert Cardiff - University of California, Davis) were cultured in DMEM supplemented with 10 % Fetal bovine serum (FBS), Penicillin/Streptomycin, L-Glucose and Sodium pyruvate (Gibco). For cell labeling with Fe₃O₄ NC, colloidosomes and supraballs, cells were incubated with the different nanoparticles at different concentrations according to the experiment for 24 hours at 37°C and 5% CO₂.

Quantification of iron uptake by cells in vitro. In order to study the uptake of the different nanoparticles in cells when incubated at [Fe] 28μ g/mL for 24 hours, the concentration of iron in A431, RAW 264.7 and hTERT-HSC cells was performed by ICP atomic emission spectrometer (model iCAP6200 duo. Thermo Fischer Scientific, analysis performed in the PARI platform, IPGP, Paris, France). For each iron measurement, cell pellets were digested in 289 μ L of 69% nitric acid (trace metal basic grade; Sigma) at room temperature overnight. The solutions were then diluted in ultrapure water for analysis. The calibration standards and the quality controls were provided by SCP SCIENCE and ChemLab. The analyzer drift is regularly corrected by quality controls measurements to ensure a drift lower than 5%. An ASX-520 Autosampler was used. Argon 4.5 Linde (min, 99.995%) was used as plasma and purge gas. All samples were analyzed in 2% nitric acid.

Fibroblast sheets culture. Cell sheets were made with hTERT-HSC, as previously described [39]. 30,000 cells were plated in 6-well plates for 5-6 weeks. During this period, the cells were supplied with DMEM freshly supplemented with 50 μ g/mL ascorbic acid (Sigma-Aldrich), three times per week. The fibroblast sheets were rinsed with PBS and incubated with the Fe₃O₄ NC, colloidosomes and supraballs at 28 μ g/mL of iron for 24 hours at 37°C and 5% CO₂. The fibroblast sheets were then washed twice with PBS and fixed with 2% glutaraldehyde in 0.1 M Na cacodylate buffer pH 7.2, for 1 hour at room temperature. Samples were then processed for Transmission Electron Microscopy as detailed in the following section.

Transmission Electron Microscopy. TEM images were acquired using a Hitachi HT7700 operating at 80 kV (MIMA2 platform, INRA, Jouy-en- Josas, France). A431 cells were exposed to Fe₃O₄ NC, colloidosomes and supraballs at 28 μ g/mL of iron for 24 hours at 37°C. Following incubation cells were detached by trypsinization, washed twice with PBS and fixed with 2% glutaraldehyde in 0.1 M Na cacodylate buffer pH 7.2, for 1 hour at room temperature. Samples were then contrasted with Oolong Tea Extract (OTE) 0.5% in cacodylate buffer for 1 hour at room temperature, postfixed with 1% osmium tetroxide containing 1.5% potassium cyanoferrate for 1 hour at room temperature, gradually dehydrated in ethanol: 70% - 5min incubation, 90% - 5 min incubation, 100% - 5min incubation 3 times. Ethanol was then substituted gradually in mix of ethanol-Epon under vacuum for 1 hour and finally embedded in Epon (Delta microscopie – Labège France).

Thin sections (70 nm) were collected onto 200 mesh copper grids, and counterstained with lead citrate. Grids were examined with Hitachi HT7700 electron microscope operated at 80kV (Elexience – France), and images were acquired with a charge-coupled device camera (AMT).

Photothermal measurement of nanoparticles in aqueous solution. Photothermal measurements were made with an 808 nm NIR laser (Laser Diode Drivers, BWT) with adjustable power setting (0-5W). 100 μ L of aqueous nanoparticle dispersion at different concentrations were placed in a 0.5 mL tube (open lid). The tube was irradiated at 4 cm distance with a laser spot of 0.4 cm². The laser power was adjusted at 1 W cm⁻². The temperature was monitored using an infrared thermal imaging camera (FLIR SC7000) in real time, recording one frame per second.

Cell necrosis assay – *global effect*. A431 cells were incubated with [Fe] $28\mu g/ml$ Fe₃O₄ NC, colloidosomes and supraballs for 24 h in 6-well cell culture plates. Cells were then trypsinized, washed with PBS1X and counted. 1x 10⁶ cells were resuspended in 100 μ l PBS 1X (500 μ l tube). The cell suspension was then subjected to an 808 nm beam laser for 10 min (1 W/cm², height 4 cm from the liquid/air interface, Laser Diode Drivers, BWT), and temperature was recorded with infrared camera SC7000 from FLIR Systems. To evaluate the global effects of heating, the irradiated cells were put back into Nunc LabTeck slides (Thermo Fisher) and cultured for 4 hours. To test the level of damage performed by the nanoparticle in cells, cell necrosis was evaluated. To do so, Chromatin Condensation & Membrane Permeability Dead Cell Apoptosis Kit, purchased from Molecular Probes-Invitrogen Detection Technologies (Eugene, OR), was used according to the manufacturer's protocol. Briefly, the cells were washed with PBS were incubated with 1 μ l each of Hoechst 33342, YO-PROTM dye and propidium iodide stock solution at 37°C in an atmosphere of 5% CO₂ for 15 minutes. The cells were viewed

using a Andor Technology with Olympus JX81/BX61 Device/Yokogawa CSU Device spinning disk microscope (Andor Technology plc, Belfast, Northern Ireland) using appropriate filters. The dead cells were stained with red-fluorescent propidium iodide dye whilst YO-PROTM dye enters cells in apoptosis, the blue-fluorescent Hoechst 33342 stains the chromatin of the cells. 30 images were taken for each condition. All experiments were performed three times. To quantify the level of cell death and apoptosis the number of propidium iodide and YO-PROTM labeled nuclei, respectively, as well as Hoechst 33342 labeled nuclei were counted using ImageJ Cell Counter plugin.

Cell necrosis assay – *local effect.* 15000 A431 cells were seeded in each well of an 18 well μ -slides (Ibidi, Germany). The cells were then incubated with Fe₃O₄ NC, colloidosomes and supraballs at 28 μ g/mL of iron overnight and irradiated to an 808 nm beam laser for 10 min (1 W/cm², height 4 cm from the liquid/air interface, Laser Diode Drivers, BWT). The cells were then incubated at 37°C, 5% CO₂ for 4 hours. Next the level of cell damage was tested using the Chromatin Condensation & Membrane Permeability Dead Cell Apoptosis Kit following the same protocol described in the previous section. The cells were viewed using a Andor Technology with Olympus JX81/BX61 Device/Yokogawa CSU Device spinning disk microscope (Andor Technology plc, Belfast, Northern Ireland) using appropriate filters. The dead cells were stained with red-fluorescent propidium iodide dye whilst YO-PROTM dye enters cells in apoptosis, the blue-fluorescent Hoechst 33342 stains the chromatin of the cells. 30 images were taken for each condition. All experiments were performed three times. To quantify the level of cell death and apoptosis the number of propidium iodide and YO-PROTM labeled nuclei, respectively, as well as Hoechst 33342 labeled nuclei were counted using ImageJ Cell Counter plugin.

PTT on sheep tendon and second harmonic generation imaging and Scanning Electron Microscopy with a Field Emission Gun (SEM-FEG). Fresh sheep tendon was obtained from our local butcher. Tendon was cut (perpendicular to the fibers' orientation) using a sharp scalpel into 1-2 cm-sized fragments, which were embedded in 5% low-gelling-temperature agarose (type VII-A; Sigma-Aldrich) prepared in PBS. Slices (100 μ m) were cut (parallel to the fibers' direction) with a vibratome (VT 1000S; Leica) in a bath of ice-cold PBS. The slices were then placed on top of 0.4- μ m organotypic culture inserts (Millicell, Millipore) in 35-mm petri dishes containing 1.1 mL RPMI 1640 without Phenol Red. [Fe] 100 μ g of Fe₃O₄ NC, colloidosomes and supraballs were deposited on the slices. The slices were then incubated for 2 hours at 37°C to allow the nanostructures penetration into the tissue. The slices were washed to remove all material that had not penetrated the slice. The slices were then irradiated with an 808 nm beam laser for 10 min (2 W/cm², height 4 cm from the slice, Laser Diode Drivers, BWT).

In order to evaluate the damage produced by the PTT on the collagen network, the slices were imaged using inverted stand Leica SP5 microscope (Leica Microsystems GmbH, Wetzlar, Germany) coupled to a femtosecond Ti:sapphire laser (Chameleon, Coherent, Saclay, France) tuned at a wavelength of 810 or 850 nm for all experiments. SHG laser exposure parameters were set in order to enable image acquisition without inducing artefactual destruction of the tissue. SHG imaging exploits the non-centrosymmetric tightly packed arrangement of amino acids in collagen that exhibit unique optical properties in order to image them. The laser beam was

circularly polarized and a Leica Microsystems HCX IRAPO 25×/0.95 W objective was used. Second harmonic generation (SHG - collagen structure) signal was detected in epi-collection through a 405/15nm bandpass filter and detected by NDD PMT detectors (Leica Microsystems) with a constant voltage supply, at constant laser excitation power, allowing direct comparison of SHG intensity values. Another way of evaluating the damage produced on the collagen network was evaluating with Scanning Electron Microscopy. After irradiation the collagen slices were directly deposited on a conductive double-faced adhesive carbon tape. The samples were carbon-coated before observations. Images were obtained with a SEM-FEG Zeiss Merlin Compact microscope operating at 10 kV (MPQ laboratory, Université de Paris).

PTT on EGI-1 tumor slices and second harmonic generation imaging. Animal experiments were performed in accordance with the French Animal Research Committee guidelines and all procedures were approved by a local ethic committee (No 14678). $2x10^6$ cells suspended in 60 µL of PBS were mixed with 60 µL of Matrigel® growth factor reduced (Corning) and implanted subcutaneously into the flank of 5-week-old female NMRI-nu (nu/nu) mice (Envigo, France). Mice were housed under standard conditions in individually ventilated cages enriched with a nesting material and kept at 22°C on a 12-h light/12-h dark cycle with ad libitum access to food and tap water. When the tumor volume reached 800-1000 mm³ (approximately 30-35 days after the injection of EGI-1 cells), tumors were considered ready for harvesting. The tumors were then resected and cut into 1-2 cm fragments. which were embedded in 5% low-gelling-temperature agarose (type VII-A; Sigma-Aldrich) prepared in PBS. Slices (100 µm) were cut (parallel to the fibers' direction) with a vibratome (VT 1000S; Leica) in a bath of ice-cold PBS. The slices were then placed on top of 0.4-µm organotypic culture inserts (Millicell, Millipore) in 35-mm petri dishes containing 1.1 mL RPMI 1640 without Phenol Red. [Fe] 100 µg of Fe₃O₄ NC, colloidosomes and supraballs were deposited on the slices. The slices were then incubated for 2 hours at 37°C to allow nanoscturctures penetration into the tissue. The slices were washed to remove all material that had not penetrated the slice. The slices were then irradiated with an 808 nm beam laser for 10 min (2 W/cm², height 4 cm from the slice, Laser Diode Drivers, BWT). In order to evaluate the damage produced by the PTT on the collagen network, the slices were imaged using inverted stand Leica SP5 microscope (Leica Microsystems GmbH, Wetzlar, Germany) coupled to a femtosecond Ti:sapphire laser (Chameleon, Coherent, Saclay, France) tuned at a wavelength of 810 or 850 nm for all experiments. The laser beam was circularly polarized and a Leica Microsystems HCX IRAPO 25×/0.95 W objective was used. Second harmonic generation (SHG - collagen structure) signal was detected in epi-collection through a 405/15nm bandpass filter and detected by NDD PMT detectors (Leica Microsystems) with a constant voltage supply, at constant laser excitation power, allowing direct comparison of SHG intensity values.

In vivo studies in MET-1 tumor models. Animal experiments were performed in accordance with the French Animal Research Committee guidelines and all procedures were approved by a local ethic committee. 1 x 10⁶ cells suspended in 50 μ L of PBS were injected subcutaneously into the flank of a 6-week-old female FVB mice (Janvier France). Tumor growth was followed with a caliper, and tumor volume (V) was calculated as follows: xenograft volume = (xy²)/2 where x is the longest and y, the shortest of two perpendicular diameters. When the

tumor volume reached 150-200 mm³ (approximately 10 days after the injection of MET-1 cells), tumors were considered ready for treatment. [Fe] = $150 \ \mu g$ of Fe₃O₄ NC, colloidosomes and supraballs were injected intra tumorally within the MET-1 tumors. 24 hours after the injection, tumors were irradiation with an 808-nm at 2 W/ cm² for 15 min (height from tumor skin 4 cm, Laser Diode Drivers, BWT). Surface temperature was recorded with an infrared camera SC7000 from FLIR Systems during the laser treatment and analyzed using ResearchIR Software (FLIR). 24 hours after irradiation the animals were sacrificed and the tumor was resected for histological analysis

Histology and immunohistochemistry. Freshly harvested tumors were fixed overnight at 4 °C in a periodate– lysine–paraformaldehyde solution [0.05 M phosphate buffer containing 0.1 M L-lysine (pH 7.4), 2 mg/mL NaIO₄, and 10 mg/mL paraformaldehyde]. The tumors were dehydrated in graded solutions of ethanol and embedded in paraffin. For histology, five micrometer tissue sections were stained with hematoxylin / eosin / Saffron (HES) and Perls staining. Slices were then imaged with an optical microscope and analyzed. For immunohistochemistry labelling, five micrometer tissue sections were dewaxed. For F4/80 labeling, the antigen was retrieved with citrate solution (pH 7) preheated for 10 min at 95-98°C. Endogenous peroxide was blocked for 10 min with 10% of hydrogen peroxide. The F4/80 antibody (D2S9R clone, Cell Signaling) was incubated at a 1/100 dilution. An automated staining system (Leica Bond) was used to perform the immunostaining. The anti-F4/80 antibody was detected with a secondary anti-rabbit antibody, coupled to a HRP-coupled polymer (Leica, Bond Polymer Refine Detection, No. DS9800).

Statistical analysis. Results were analyzed using the GraphPad Prism 5.0 statistical software. Data are shown as means \pm standard error of the mean (SEM). For comparisons between two groups, parametric Student t-test or non-parametric Mann-Whitney test were used. For comparisons between more than two groups, parametric One-Way analysis of variance (ANOVA) test followed by a posteriori Kruskal-Walis test.

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Conflict of Interest

The authors declare no conflict of interest.

Keywords

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