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Computationally Guided Photothermal Tumor Therapy Using Long-Circulating Gold Nanorod Antennas

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Abstract

Plasmonic nanomaterials have the opportunity to considerably improve the specificity of cancer ablation by i.v. homing to tumors and acting as antennas for accepting externally applied energy. Here, we describe an integrated approach to improved plasmonic therapy composed of multimodal nanomaterial optimization and computational irradiation protocol development. We synthesized polyethylene glycol (PEG)-protected gold nanorods (NR) that exhibit superior spectral bandwidth, photothermal heat generation per gram of gold, and circulation half-life *in vivo* ($t_{1/2}$, ~17 hours) compared with the prototypical tunable plasmonic particles, gold nanoshells, as well as ~2-fold higher X-ray absorption than a clinical iodine contrast agent. After intratumoral or i.v. administration, we fuse PEG-NR biodistribution data derived via noninvasive X-ray computed tomography or *ex vivo* spectrometry, respectively, with four-dimensional computational heat transport modeling to predict photothermal heating during irradiation. In computationally driven pilot therapeutic studies, we show that a single i.v. injection of PEG-NRs enabled destruction of all irradiated human xenograft tumors in mice. These studies highlight the potential of integrating computational therapy design with nanotherapeutic development for ultraselective tumor ablation.

Introduction

The electromagnetic properties of plasmonic nanomaterials have been harnessed to develop ultrasensitive diagnostic (1,2), spectroscopic (3,4), and, recently, therapeutic technologies (5-8). In particular, tunable plasmonic nanomaterials have attracted attention for their immense optical absorption coefficients and potential as injectable nanoantennas that target tumors and locally convert otherwise benign electromagnetic energy to thermal energy for ablation. Currently, tumor ablation approaches in clinical practice, including radio frequency, laser, and

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focused ultrasound methods, lack intrinsic tumor specificity to energy absorption. The inability to selectively heat tumor tissues over surrounding compartments necessitates efforts to externally direct applied energy toward tumor tissues, making effective treatment of tumor margins and complex tumor geometries very challenging. By providing a tumor-specific heat source, nanoantennas could considerably broaden the clinical applicability of thermal therapies by simplifying their integration with current therapeutic practices (including improving margin clearance in surgery and synergizing with regional radiation therapies) and reducing morbidity due to off-target heating. Furthermore, by pulsing the external energy source used, tumor-targeted nanoantennas can theoretically ablate with single-cell precision, thereby providing improved accuracy over standard surgical methods and opening the possibility of precisely treating complex tumor margins in sensitive tissues.

To date, preparations of gold nanoshells and nanorods (NR) have shown considerable efficacy for tumor ablation using NIR light (5,6,9,10), with the most recent data showing complete resorption of ~55% and ~25% of irradiated tumors, respectively (11,12). These results highlight the clinical promise of these technologies and also motivate the further development of superior nanomaterials and improved methods for optimizing irradiation regimens, which could synergistically improve photothermal therapies. From a material perspective, the development of nanoantennas with enhanced circulation times in vivo, increased absorption coefficients per weight, and narrower absorption spectra would more efficiently permeate into tumors after i.v. administration, amplify the photothermal contrast between antennas and normal tissue, and allow improved tumor treatment at lower laser intensities or at greater depths in vivo. From a procedural perspective, despite a rich history of heat transfer modeling in the hyperthermia field, the use of quantitative modeling to predict the in vivo function of plasmonic nanomaterials has widely remained absent from their testing (5,6,9,11,12). Because the efficacy of photothermal therapy is driven by both the potency of nanoantenna absorption in tumors and the dose of near-IR irradiation, translation of plasmonic materials to effective clinical use will benefit from cohesive integration of biodistribution data with photothermal modeling to predict and customize the four-dimensional irradiated temperature profiles in tumors.

Recently, rod-shaped gold nanoparticles have emerged as precisely tunable plasmonic nanomaterials that may be synthesized in bulk, have narrow size distributions, optical absorption coefficients 10^4 -fold to 10^6 -fold higher than conventional organic fluorochromes, and theoretical per micron absorption coefficients exceeding those of NIR-absorbing gold nanoshells (13-15). The long precedence of gold nanoparticles in clinical rheumatoid arthritis therapies make gold NRs appealing new candidates for nanoantenna-based photothermal ablation and a wide array of other biomedical applications. Already, gold NRs have been used for a diversity of biological purposes, including multiplexed *in vitro* detection (16), two-photon fluorescence imaging (17), and photothermal heating of tumor and bacterial cell targets (7,8, 12,18-20). In addition to their plasmon resonance, the larger atomic number and high material density of gold nanomaterials (z = 79, $\rho = 19.3$ g/cm³) compared with clinical formulations of iodine-based reagents (z = 53) have attracted interest for X-ray computed tomography (CT) angiography and a few spherical nanoparticle reagents have been developed for *in vivo* use (21,22).

In this report, we describe the development of polyethylene glycol (PEG)-coated gold NRs as highly absorbing nanoantennas for photothermal tumor destruction under the guidance of biodistribution-based photothermal modeling. We chose a PEG polymer coating due to the widespread clinical use of variable-length PEG polymers for extending the circulation time of protein therapeutics (23,24) and for nanoparticle formulations, such as the drug-loaded liposomes Doxil. We find that PEG-NRs are highly stable, relatively noncytotoxic *in vitro*, superior photothermal antennas compared with a gold nanoshell preparation *in vitro* and are improved X-ray absorbing agents compared with clinical iodine standard. After i.v.

administration, we find PEG-NRs to be among the longest circulating inorganic nanomaterials described to date ($t_{1/2}$, ~17 hours) allowing passive accumulation into xenograft tumors. Using four-dimensional biodistribution-based heat transfer simulations, we designed a therapeutic irradiation regimen that was able to fully destroy all irradiated tumors on mice injected with PEG-NRs using half the light intensity of previous nanoshell therapies.

Materials and Methods

Preparation of PEG-coated gold NRs

Highly stable, ~13 × 47 nm (Fig. 1A) cetyltrimethylammonium bromide (CTAB)-coated gold NRs with longitudinal plasmon resonance at 810 nm (Nanopartz, a division of Concurrent Analytical, Inc.) were centrifuged at 16,000 rcf to concentrate and gently resuspended in 250 µmol/L 5-kDa methyl-PEG-thiol (Laysan Bio, Inc.). Thiol activity of polymers was quantified spectraphotometrically using 5,5-dithiobis(2-nitrobenzoic acid) (Sigma) against a DTT (Sigma) gradient to verify that polymer stocks were fully reduced. The solution of 5-kDa methyl-PEG-thiol and CTAB-coated gold NRs was gently mixed at room temperature for 1 h and dialyzed exhaustively against ultrapure water (18 M Ω cm⁻¹) via cellulose ester membrane dialysis to drive PEG addition (Spectrapor). Dialyzed samples were filtered through 100-kDa filters (Millipore) to remove excess polymer and stored at 4jC. To quantify the number of polymers per particle, NRs were coated as described with an amino-PEG-thiol polymer. After dialysis and extensive filtration on 100-kDa centrifugal filters, amino NRs were harvested and an SPDP assay was performed to quantify the number of amines (25).

Stability and cytotoxicity

Solutions of PEG-NRs or CTAB-NRs (\sim 60 µg Au/mL) were normalized and incubated in PBS or 10% human serum for extended periods of time. At regular intervals, samples were spectrophotometrically analyzed for plasmon resonance peak shifts, which would indicate particle aggregation. To assess material toxicity, micropatterned primary hepatocyte/human fibroblast cocultures were prepared as described previously (26). At 24 h after liver coculture seeding, coculture wells were exposed in triplicate to a gradient of PEG-NRs (0-280 µg Au/mL) and allowed to incubate for 24 h. At this point, PEG-NRs were removed, cells were washed repeatedly, and viability was assessed via a 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide viability assay (Sigma), a colorimetric viability assay for mitochondrial dehydrogenase enzyme activity read using a spectrophotometer (SpectraMax, Molecular Devices).

Photothermal comparison between PEG-NRs and PEG-nanoshells

Gold nanoshells (Nanocomposix) with 120 nm silica core and ~15 nm gold shell were mixed with methyl-PEG-thiol as described (5,6,9). Both PEGnanoshells and PEG-NRs were brought to 7 μ g Au/mL, as determined by inductively coupled plasma mass spectrometry (ICP-MS). Tubes containing 200 μ L of these solutions were broadly irradiated, under identical conditions, by an 810-nm diode laser (RPMC Lasers, Inc.) at 2 W/cm². During irradiation, an IR thermographic camera (FLIR, Thermacam S60) was used to measure peak sample surface temperature. To assess photothermal cell toxicity, MDA-MB-435 human cancer cells [American Type Culture Collection (ATCC)] were cultured in a 96-well microplate and grown to 80% confluency using ATCC-recommended media. Cells were incubated with either PEG-NRs (14 μ g/mL), PEG-nanoshells (14 Ag/mL), or media alone. For each, triplicate wells were exposed to the diode laser light (5 min, 2 W/cm²) or no laser. After treatment, cells were incubated with Calcein AM (5 Ag/mL in culture medium; 1 h incubation, Invitrogen), a fluorescent indicator of esterase activity in viable cells and imaged using phase and fluorescence microscopy.

X-ray CT of PEG-NRs

PEG-NRs were suspended in PBS after concentration via membrane centrifugation and serially diluted over a 1,000-fold concentration gradient. A clinical iodine standard was similarly diluted for comparison (Isovue-370). X-ray CT was performed in a GE eXplore Locus microCT scanner (80 kV, 450 μ A, 45- μ m resolution). For *in vivo* imaging, mice were imaged before NR injection to reveal the baseline level of soft tissue X-ray contrast. Ten microliters of PEG-NRs (~3 pmol) were injected with a 30-gauge needle directly into the center of the tumor, and the needle was maintained in place for ~10 s to allow the tumor to accommodate the additional fluid. After intratumoral administration, mice were imaged and irradiated (~0.75 W/cm², 810 nm). X-ray CT images were exported as DICOM files for exportation into modeling software (see Supporting Materials and Methods).

ICP-MS

Samples for ICP-MS (Thermo-Scientific Finnigan ELEMENT2) analysis were frozen, lyophilized, and dissolved in aqua regia, prepared by adding 100 μL of nitric acid + 300 μL of 37% hydrochloric acid for 72 h to dissolve gold particles. Then, samples were diluted to 10 mL with 9.6-mL 2% HNO3 and analyzed via ICP-MS against standards. Control saline and organ samples with exogenously added PEG-NRs were used to calibrate this method.

In vivo circulation time and biodistribution of PEG-NRs

Nude mice were bilaterally injected s.c. in the hind flank with ~ 2×10^6 MDA-MB-435 cells. After 2 to 3 wk, animals were anaesthetized with isoflurane and injected through the tail vein with PEG-NRs in 0.15 mol/L NaCl, 0.1 mol/L Na phosphate buffer (pH 7.2; 20 mg Au/kg). 10 μ L blood samples were withdrawn periodically from the suborbital space, diluted with PBS containing 10 mmol/L EDTA, centrifuged to remove RBCs, and read on a spectrophotometer to assess PEG-NR plasmon peak height. For biodistribution experiments, after vascular clearance of PEG-NRs (72 h), injected animals were euthanized and organs were collected, weighed, and lyophilized for ICP-MS quantification of PEG-NR biodistribution.

In vivo photothermal heating of gold NRs and photothermal therapy

For both initial modeling and growth curve assessments after photothermal treatment, nude mice were bilaterally injected in the hind flank with $\sim 2 \times 10^6$ MDA-MB-435 cells. After 2 to 3 wk, animals were anaesthetized with isoflurane and injected through the tail vein with PEG-NRs in PBS (20 mg Au/kg) or PBS alone. After vascular clearance of PEG-NRs (72 h), the right flank of each mouse was irradiated (2 W/cm², 810 nm, 1 cm beam diameter). Thermographic imaging of photothermal heating was used to facilitate modeling of threedimensional temperature distributions in tumors (n = 3 mice for each set). To explore the hematologic effects of NR administration and near-IR ablation, mice bearing bilateral MDA-MB-435 tumors were injected with saline or PEG-NRs and, 72 h later, either exposed to the therapeutic tumor irradiation protocol under anesthesia (~2 W/cm², 5 min, 810 nm) or anesthesia without irradiation (n = 3 each set). After exposure, blood was collected for hematology and mice were sacrificed. Therapeutic assessment of the affect of PEG-NR heating on tumor growth was conducted similarly (n = 4 mice in each treatment set). Both irradiated and unirradiated tumors of each mouse in the therapeutic assessment trial were measured at regular intervals using digital calipers. Survival studies were conducted using mice that were unilaterally injected in the hind flank with $\sim 2 \times 10^6$ MDA-MB-435 cells (n = 5 mice in each treatment set). Tumor sizes were measured over time, and mice were euthanized once tumors exceeded 500 mm³.

Results

Development of ultrastable, polymer-coated gold NRs

In principle, gold nanoparticles are attractive for medical applications because various formulations have been approved by the U.S. Food and Drug Administration and in clinical use for decades. However, one barrier facing the widespread biological use of gold NRs has been the need to replace the cationic CTAB surfactants used to drive their anisotropic growth with hydrophilic, biocompatible coatings. We found that NRs, with axial sizes of 12.7 nm (± 3.4) and 47 nm $(\pm 9.3;$ Fig. 1A), coated in CTAB are cytotoxic and colloidally unstable in 0.15 mol/L NaCl or 10% human serum (Fig. 1B and C). After PEGylation (Fig. 1B), gold NRs contained ~20,000 polymers per particle by the SPDP assay (25) and were rendered highly stable in vitro, showing minimal spectral shifting (which would indicate particle destabilization and aggregation) even after >1,000 hours in 0.15 mol/L NaCl or 10% human serum (Fig. 1C and Supplementary Fig. S1A). Additionally, PEG-NRs could be dispersed in a variety of solvents, including acetone, acetonitrile, DMSO, DMF, ethanol, and methanol, further highlighting the stability of their coating and their amenability to future chemical processing and functionalization (Supplementary Fig. S1B). After 24 hours of incubation above primary rat hepatocyte cocultures, an in vitro liver model that was previously used to elucidate semiconductor quantum dot toxicity (26) and has shown utility for rank-ordering pharmacologic toxicities (27), PEG-NRs, displayed no significant alterations in mitochondrial activity compared with saline alone, even at doses 20 times greater than that used over cells in vitro here and equal to maximal blood concentrations during in vivo therapy experiments (Supplementary Fig. S2), highlighting their potential biocompatibility for medical applications.

Photothermal comparison of gold NRs and gold nanoshells

In light of the prior utility of gold nanoshells for photothermal tumor therapy, they were used as a benchmark here to evaluate the photothermal performance of PEG-NRs (Fig. 2A): PEGylated nanoshell preparations with similar composition and spectra to those used for photothermal *in vivo* applications (refs. 5,6,9; ~120 nm silica cores and ~15 nm gold shells, 810 nm peak plasmon resonance; Fig. 2B). PEG-NRs exhibited <1/3 of the spectral bandwidth and ~3 times higher extinction coefficient per gram gold than PEG-nanoshells (full width at half maximum of PEG-NRs 130 nm, PEG-nanoshells 490 nm; Fig. 2B). Additionally, under identical experimental conditions, irradiated PEG-NR solutions generated heat >6 times more rapidly than PEGnanoshells per gram gold (Fig. 2C). These findings are consistent with theoretical calculations indicating that gold NRs of this size exhibit superior per micron absorption coefficients to gold nanoshells (14). Accordingly, incubation of PEG-NRs with MDAMB-435 human tumor cells *in vitro* enabled their widespread destruction with levels of NIR light that were insufficient for nanoshell-mediated toxicity (Fig. 2D). Exposure to individual stimuli (NRs, nanoshells, or laser) did not affect cell viability (Supplementary Fig. S3).

X-ray CT and computational modeling of photothermal NR heating in vivo

To translate the photothermal efficacy of PEG-NRs to *in vivo* therapy, we next developed a method through which PEG-NRs could be heated under the guidance of biodistributionbased computational photothermal modeling. To model and customize patient irradiation regimens, we sought to acquire nanoparticle distributions remotely using X-ray CT, a desirable imaging modality due to its high three-dimensional anatomic resolution, rapid imaging speed, quantitative dynamic range of detection, full body penetration, and ubiquitous clinical use. To investigate the ability of PEG-NRs to act as dense X-ray absorbing agents for X-ray CT, solutions of PEG-NRs were serially diluted and imaged using a GEmicroCT scanner (Fig. 3A). The resulting relationship between PEG-NR concentration and X-ray contrast was highly

linear and exhibited ~2-fold amplified X-ray contrast compared with a clinical iodine standard per mole (Fig. 3*B* and Supplementary Fig. S4), analogous to that found previously for spherical gold nanoparticle reagents (21,22). In addition to providing enhanced X-ray absorption, PEG-NRs allow NIR photothermal heating whereas iodine reagents show no heating above water alone (Supplementary Fig. S4).

To assess the utility of the high X-ray absorption of PEG-NRs for remote detection in vivo, ~3 pmol of PEG-NRs (~1 µmol Au) were directly injected into the tumors of mice bearing bilateral human MDA-MB-435 tumors, implanted either in the mammary fat pad or the rear flank. We found that X-ray CT rapidly detailed the threedimensional distribution of PEG-NRs in tumors, showing clear distinction between NRs and soft tissues (Fig. 3C and Supplementary Figs. S5 and S6). To understand the relationship between the nanoparticle distribution in tumors and the associated processes of thermal deposition and flux during irradiation, a finite element model of PEG-NR heating was developed based on the Pennes bioheat transfer equation, including (a) three-dimensional skeletal, soft tissue, and PEG-NR geometries transferred from X-ray CT imaging; (b) temperature-dependent optical absorption and scattering; (c) heat transfer, including surface thermal flux, internal diffusive flux, and temperature-dependent perfusive thermal flux in tissues; and (d) an extracorporeal NIR laser of variable intensity, beam diameter, and orientation matching used (see supplementary data). X-ray CT data was exported into ScanIP and ScanFEimage processing software for skeletal, NR, and soft tissue geometry construction and subsequently into COMSOL Multiphysics modeling software for photothermal simulations. Exported geometries enabled rapid delineation PEG-NRs, along with skeletal structures and surrounding soft tissues for spatially defining model parameters (Fig. 3C and Supplementary Figs. S5 and S6). Computational, finite element heat transfer simulations were performed using PEG-NRs and tumor geometries as computational domains to evaluate the feasibility of applying four-dimensional modeling to the process of photothermal heating under irradiation (Fig. 3D and Supplementary Fig. S6). Simulations revealed the intratumoral vectors of thermal flux extending from regions of PEG-NRs, as well as the internal and surface temperature history expected for irradiation at varying laser intensities (Fig. 3D and Supplementary Fig. S6). At matched computational and experimental laser intensities, simulated surface heating behavior qualitatively and quantitatively matched the observed surface temperature values and distributions acquired using IR thermographic observation (Fig. 3D and Supplementary Fig. S6). We believe this establishes the potential of fusing quantitative imaging with computational modeling to provide insight into the unintuitive, highly complex phenomena of in vivo photothermal heating. Next, we proceeded to explore the power of this modeling to quantitatively predict in vivo heating after i.v. tumor targeting.

Long circulation time and photothermal tumor heating after i.v. NR administration

We next investigated the behavior of PEG-NRs after i.v. administration in mice to systemically target tumors through the enhanced permeability and retention effect (28). For PEG-NRs to passively target tumors via the enhanced permeability and retention effect and act as nanoantennas for photothermal therapy, they must be able to traverse the systemic circulation, deter protein opsonization and reticuloendothelial system (RES) clearance, permeate through transendothelial pores in tumor blood vessels, and be retained in the tumor interstitium. After i.v. administration to tumor-bearing mice (20 mg Au/kg), our PEG-NRs were found to exhibit blood half-lives of f17 hours (Fig. 4A) and to maintain their 810-nm longitudinal plasmon resonance throughout this time, allowing spectrophotometric detection in serum over time (Supplementary Fig. S7).

To quantitatively assess tumor accumulation of PEG-NRs, nude mice bearing MDA-MB-435 human tumors were given i.v. PEG-NRs, and once NRs had cleared from circulation (72 hours),

organs were removed for gold quantitation. Here, ICP-MS was used to quantify the accumulation of exogenously given gold in tissues. ICP-MS Au NR detection was found to be highly sensitive and linear across a relevant range for gold NR detection (Supplementary Fig. S8). As expected for nanomaterials above the renal filtration limit, PEG-NRs were found to clear via RES uptake with splenic clearance dominating hepatic (Fig. 4B; Supplementary Table S1), a pattern analogous to that observed previously for PEG-protected 10-nm diameter spherical gold nanoparticles (21). Importantly, passive tumor accumulation of PEG-NRs after injection was found to be highly efficient, even at 72 hours after injection, accumulating at ~7% ID/g (Fig. 4B; Supplementary Table S1), allowing PEG-NRs to amplify the tumor absorption coefficient of 810 nm light by ~7 fold (see supplementary data). Based on empirical studies of passive tumor targeting, the enhancement of the PEG-NR circulation time over reported PEG-nanoshell formulations should amplify their passive tumor accumulation in tumors, which, in concert with their enhanced photothermal heat generation in vitro, would be expected to provide overall enhanced photothermal contrast between tumors and normal tissue. However, as PEGnanoshells are not commercially available, a side-by-side in vivo comparison could not be pursued. Separately, to study the long-term clearance of PEG-NRs from RES organs, tumor-free mice were injected with PEG-NRs and sacrificed 2 months after injection. During these 2 months, injected mice showed no signs over NR toxicity, such as weakness, belabored breathing, or failure to thrive. Organ analysis showed that the %ID/g values for tissues decreased by >50% in all organs and by >80% in muscle, heart, lungs, and kidneys (Supplementary Fig. S9), indicating the existence of routes for gold NR clearance after uptake in tissues.

Having observed their efficient passive homing to human tumors via the enhanced permeability and retention effect, the ability of PEG-NRs to remain active as optical nanoantennas for photothermal tumor heating after clearance from the systemic circulation (~72 hours) was subsequently investigated. Either PEG-NRs (20 mg Au/kg) or a saline solution was given i.v. into mice bearing two MDA-MB-435 tumors on opposite flanks. Once PEG-NRs had cleared from circulation (72 hours), irradiated tumors on PEG-NR-injected mice rapidly heated to temperatures of over 70°C (Fig. 4C), whereas saline-injected mice displayed less focal temperature increases with maximum surface temperatures of ~40°C(Fig. 4D). To inform the development of near-IR radiation doses that would fully destroy tumors, photothermal heating simulations were again conducted using the i.v. biodistribution data and computational domains containing ellipsoidal tumor geometries matching those used in experiments (Supplementary Fig. S10). Photothermal heating simulations closely matched experimental surface temperature data (Figs. 4D and 5A and B), suggesting that tumor-targeted nanoantennas substantially retained their photothermal efficacy during the 72 hours after injection in vivo. Furthermore, the simulations predicted that the entire tumor volume, including the deepest tumor/tissue interface, would be heated to ablative temperatures (>60°C) by 5 minutes after the onset of laser irradiation (Fig. 5C). Thus, a 5-minute, 2-W/cm² irradiation regimen was used for subsequent therapeutic experiments in an effort to provide fully destructive photothermal tumor treatment.

Photothermal tumor destruction using a computationally designed irradiation regimen

To test the prediction that a single dose of PEG-NRs could destroy tumors under the computationally designed protocol of NIR irradiation, nude mice bearing bilateral human MDA-MB-435 tumors were injected with either PEG-NRs or saline. After i.v. clearance of PEG-NRs, the right flank of each mouse was irradiated for 5 min (810 nm, 2 W/cm²) and all tumors were measured at regular intervals over time. Within 10 days all the irradiated, PEG-NR-targeted tumors completely disappeared by external observation whereas all other tumors, including those exposed to laser after saline injection, continued to grow uninhibited (Fig. 6A). To assess the survival benefit of PEG-NR-directed tumor ablation, mice bearing a single

MDA-MB-435 tumor were divided between four cohorts (PEG-NRs + laser, PEG-NRs - laser, saline + laser, saline - laser) and all tumors were measured over time (Fig. 6B). By 20 days after treatment, all irradiated, PEGNR-injected mice displayed only a minor scar with no evidence of tumor regrowth, whereas all other surviving mice harbored thriving tumors (Fig. 6B and C). Over the course of >50 days of observation, no irradiated, PEG-NR-injected mice showed evidence of recurrence whereas all mice in the control had to be euthanized by day 33. Body weights of PEG-NR-treated mice were monitored over time and showed no obvious losses due to tumor ablation and resorption (Supplementary Fig. S11). In a separate experiment to assess the acute hematologic effects of NR-directed tumor ablation, the only statistically significant change observed in response to NR-mediated tumor ablation was a slight increase in the percentage of band neutrophils in NR + laser sets (P < 0.05 for NR+ laser versus NR, saline + laser, and saline; Supplementary Fig. S12), likely due to an acute inflammatory response to tumor ablation.

Discussion

Here, we present the development of an integrated system for nanoantenna-based photothermal tumor therapy involving the synthesis of long-circulating gold NRs as efficient NIR-nanoantennas, biodistribution data acquisition via X-ray CT nano-material imaging or *ex vivo* spectrometry, and photothermal computational modeling to guide surgical irradiation planning. Broadly, the efficacy of a nanoantenna for photothermal therapy depends on both intrinsic (optical absorption coefficient and material cytotoxicity) and extrinsic (polymer coating, macrophage affinity, and circulation time) material characteristics, as well as external parameters, such as the use of optimized dosing and irradiation protocols for effective treatment.

We show that PEG-NRs exhibit superior intrinsic absorption and photothermal efficacy compared with gold nanoshells (~6 times greater heat generation per weight gold), as well as substantially improved circulation times *in vivo* (~17 hours versus ~4 hours), extrinsically imparted by their polymer coating (5,6). Surveying literature on inorganic nanoparticle circulation times *in vivo*, the circulation half-life of PEG-NRs is among the longest achieved to date. Previously, polymer-stabilized inorganic nanomaterials have been described with circulation half-lives of a few hours *in vivo* (29,30), including a variety of other gold nanoparticle preparations (22,31-33), and on occasion with circulation times of ~10 to 15 hours in mice (21,28,34). Elsewhere, another PEG-NR formulation was developed for *in vivo* applications, but showed a 30-minute half-life without investigation into their ability to passively target tumors or mediate *in vivo* photothermal heating (32). Because nanoparticle circulation time has been shown to determine the efficiency of nanoparticle accumulation in tumors via the enhanced permeability and retention effect, in mouse cancer models and clinical cancer treatment (28), the long circulation time reported here has the potential to directly translate to improved clinical tumor accumulation over previous nanoantennas.

Beyond the material determinants of nanoantenna efficacy, the irradiation protocol used (i.e., beam intensity, shape, cross-section, duration, direction, etc.) and nanoantenna dosing regimen directly control the rates of energy capture and dissipation to surrounding tissues *in vivo*. Whereas nanoantennas have the potential to increase the selectivity of tumor ablation, unoptimized irradiation of tissues carries the risks of either unnecessary morbidity due to collateral damage or ineffective therapy due to incomplete treatment of tumor margins. Here, we show that quantitative biodistribution data incorporated into computational modeling can help anticipate the photothermal heating in tumors and surrounding tissues during irradiation. Future developments of the quantitative model presented here could enable rapid quantitative modeling of photothermal temperature gradients in arbitrarily complex three-dimensional tissues and provide a route toward *a priori* personalization of irradiation regimens. As a proof

of principle, we establish a means of integrating whole-subject X-ray CT data with quantitative heat transfer modeling, offering a new route toward merging the clinical paradigms of imaging and therapy for personalized four-dimensional radiation planning and optimization. Furthermore, using a computationally planned therapeutic method, we show that i.v. administration of PEG-NR nanoantennas enabled complete destruction of all irradiated tumors under otherwise benign near-IR light.

We believe our findings motivate future investigation into the long-term biodistribution of PEG-NRs, more extensive analysis of their potential toxicity *in vivo*, and the development of methods for detecting low concentrations of PEG-NRs in whole animals to remotely quantify i.v. tumor targeting. Methods for actively targeting NRs to tumors, particularly to vascular epitopes, could potentially enhance their specificity for tumors or direct their additional accumulation in premalignant lesions and metastatic lymphatics. Finally, we provide clear evidence that the application of quantitative biodistribution-based modeling to the *in vivo* testing of nanomaterials can provide insight into their function and direct procedural optimization.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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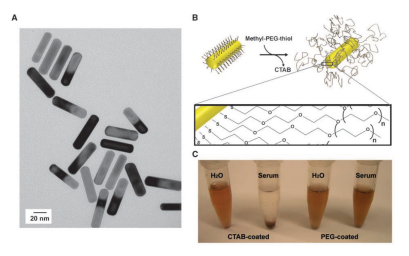


Figure 1. Structure and synthesis of highly absorbing, PEG-protected gold NRs. *A*, near-IR absorbing (810nm longitudinal plasmon resonance peak) gold NRs were imaged via transmission electron microscopy. *B*, schematic of process to drive CTAB-NR conversion to PEG-NRs under dialysis with rendering and molecular schematic of PEG coating on NR surface. *C*, PEG-NRs show prolonged stability in biological media (>1,000 h), whereas CTAB-coated NRs precipitated over time.

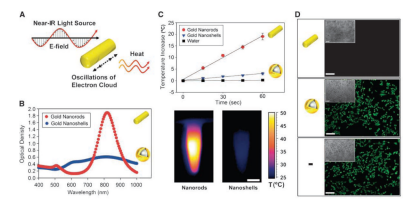


Figure 2. Spectral and photothermal properties of highly absorbing gold NRs compared with gold nanoshells. *A*, schematic of photothermal heating of gold NRs. The dimensions of gold NRs are tuned to have a near-IR plasmon resonance, at which point nanoparticle electrons resonantly oscillate and dissipate energy as heat. *B*, spectra for PEG-gold NRs (red) and PEG-gold nanoshells (blue), a benchmark for tunable plasmonic nanomaterials, at equal gold concentrations. *C*, top, rate of temperature increase for triplicate PEG-NR and PEG-gold nanoshell solutions (7 μg Au/mL, 810nm laser, 2 W/cm², n = 3 each). Bottom, IR thermographic image of PEG-NRs versus PEG-gold nanoshells after 2 min of irradiation. Scale bar, 5 mm. *D, In vitro* photothermal toxicity of PEG-NRs over human cancer cells in culture (MDA-MB-435). Tumor cells were incubated with PEG-NRs (14 μg/mL; top), PEG-nanoshells (14 μg/mL; middle), or media alone (bottom) and treated with laser irradiation (2 W/cm², 810nm, 5 min). Calcein AM staining indicates destruction of cells with PEG-NRs, whereas cells irradiated in the presence of nanoshells or media remained viable. Phase region of calcein staining inset. Scale bar, 10 μm.

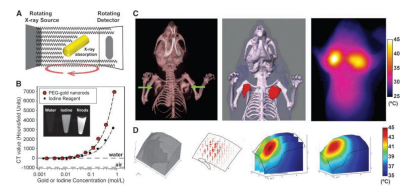


Figure 3.X-ray CT, quantitative photothermal modeling, and near-IR photothermal heating of gold NRs *In vivo. A*, schematic of X-ray absorption by gold NRs in X-ray CT. *B*, X-ray CT number of PEG-NRs compared with an iodine standard (Isovue-370). *C*, PEG-NRs were intratumorally given to mice bearing bilateral MDA-MB-435 tumors and imaged using X-ray CT to visualize three-dimensional PEG-NR distribution in tumors (left). A three-dimensional solid model of the complete geometry was rapidly reconstructed by image processing for use with computational photothermal modeling (middle). Red, PEG-NRs. Experimental thermographic surveillance of NIR irradiation after X-ray CT (~0.75 W/cm², 1 min; right). *D*, meshed geometry of the left tumor chosen as the computational domain (left). Plot of theoretical heat flux propagation inside the tumor upon irradiation (middle left). Predicted internal temperature distribution at three different planes inside the tumor (middle right) along with surface temperature map (right) matching the left tumor in C.

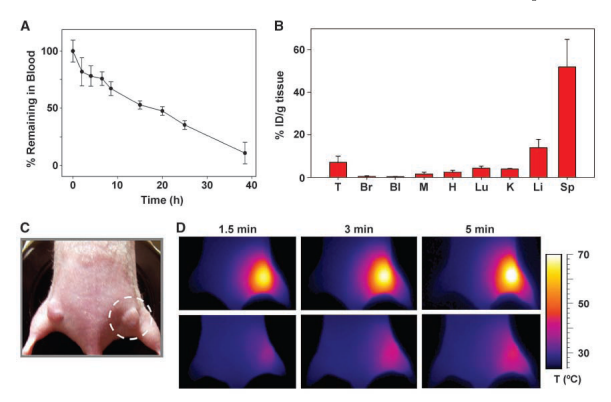


Figure 4.
Long circulation time, passive tumor targeting, and photothermal heating of passively targeted gold NR antennas in tumors. *A*, PEG-NRs were i.v. given (20mg/kg) to three mice bearing MDA-MB-435 tumors, and blood was withdrawn over time to monitor clearance from circulation. *B*, PEG-NR biodistribution and targeting to MDA-MB-435 tumors 72 h after i.v. administration, quantified via ICP-MS (three mice). T, tumor; Br, brain; Bl, bladder; M, muscle; H, heart; Lu, lung; K, kidney; Li, liver; SP, spleen. Data are tabulated in Supplementary Table S1. *C*, PEG-NRs or saline were i.v. given (20mg/kg) to mice bearing MDA-MB-435 tumors on opposing flanks. After NRs had cleared from circulation (72 h after injection), the right flank was irradiated using an 810-nm diode laser (2 W/cm²; beam size indicated by dotted circle). *D*, thermographic surveillance of photothermal heating in PEG-NR-injected (top) and saline-injected (bottom) mice.

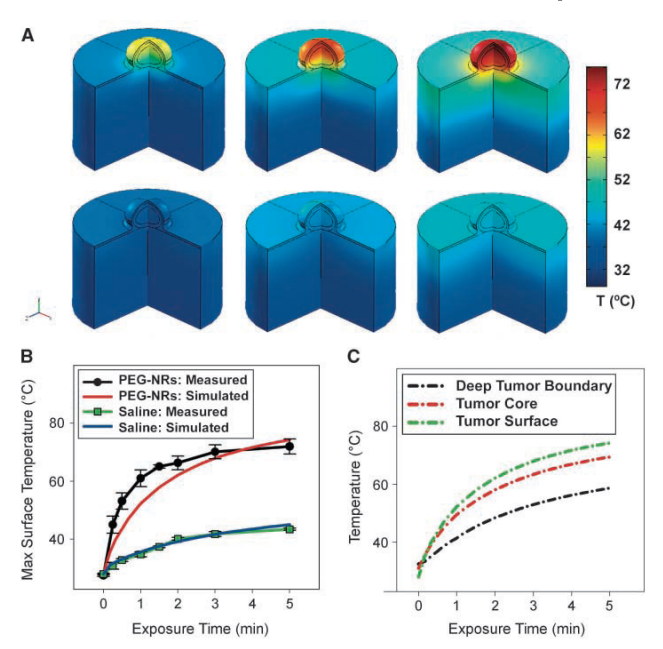


Figure 5. Quantitative photothermal modeling of gold NR tumor heating. A, three-dimensional finite element modeling of PEG-NR heating $In\ vivo$. Simulated three-dimensional temperature distributions matching the four-dimensional thermographic time points for PEG-NR (top) and control tumor irradiation (bottom). B, thermographically measured and simulated tumor surface temperatures over time for irradiation of PEG-NR or saline mice. C, simulated temperature increases various depths for PEG-NR-injected and saline-injected mice. By 5 min after the onset of irradiation, the entire PEG-NR tumor is predicted to have reached ablative temperatures of $>60^{\circ}$, motivating the choice of this irradiation regimen for subsequent therapeutic trials.

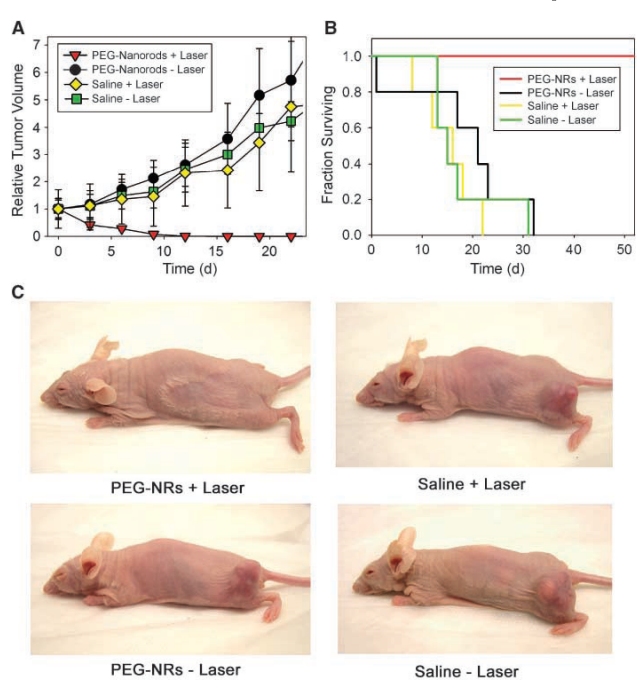


Figure 6.

Photothermal destruction of human tumors in mice using long-circulating gold NRs. *A*, mice harboring two MDA-MB-435 human tumors on opposite flanks were injected with either saline or PEG-NRs. After PEG-NRs had cleared from circulation (72 h after injection), the right flank of each mouse was exposed to the computationally designed irradiation regimen (810nm, 2 W/cm², 5 min). Volumetric changes in tumor sizes are plotted over time after irradiation. *B*, mice harboring one MDA-MB-435 human tumor were injected with either saline or PEG-NRs and irradiated as in A. Survival of mice after irradiation is plotted versus time after irradiation. *C*, at 20d after irradiation, NIR-irradiated, all PEG-NR-injected mice showed only a minor scar

and no evidence of tumor regrowth whereas all other treatment groups harbored thriving tumors.