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Enhanced detection of early-stage oral cancer in vivo by optical coherence tomography using multimodal delivery of gold nanoparticles

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Abstract

Contrast in optical coherence tomography (OCT) images can be enhanced by utilizing surface plasmon resonant gold nanoparticles. To improve the poor in vivo transport of gold nanoparticles through biological barriers, an efficient delivery strategy is needed. In this study, the improved penetration and distribution of gold nanoparticles were achieved by microneedle and ultrasound, respectively, and it was demonstrated that this multimodal delivery of antibody-conjugated PEGylated gold nanoparticles enhanced the contrast in in vivo OCT images of oral dysplasia in a hamster model.

Keywords

optical coherence tomography; optical contrast agent; gold nanoparticles; enhanced delivery; ultrasound; microneedles

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1 Introduction

The sensitive and specific detection of premalignant and malignant lesions is a crucial and indispensable prerequisite for optimal treatment outcome and prognosis. Surgical biopsy, which is currently the primary diagnostic tool, often causes patient anxiety and discomfort, undesirable side effects, and sampling errors. For example, multiple prostate biopsies miss up to 30% of cancer present at the time of biopsy.\textsuperscript{1–3} An urgent need exists to develop cost-effective, minimally invasive imaging technologies to identify premalignancy and malignancy with high sensitivity and specificity.\textsuperscript{4} Optical imaging tools such as reflectance confocal microscopy, optical coherence tomography (OCT), and optical coherence microscopy (OCM) are promising diagnostic tools for detecting cancer at an early stage.\textsuperscript{5} Optical imaging is particularly useful for \textit{in vivo} applications because it uses near-infrared wavelengths (770 to 1300 nm) that avoid predominant absorption in tissue.\textsuperscript{6}

OCT, which detects a reflected light source from the sample using refractive index mismatching, is particularly promising for early diagnosis of neoplasia because it provides resolution that is an order of magnitude higher than that of other minimally invasive diagnostic techniques such as computed tomography (CT), magnetic resonance imaging (MRI), and ultrasound (US).\textsuperscript{6,7} Real-time OCT imaging with ultrafast image acquisition can minimize the errors generated by patient and operator motion artifacts.\textsuperscript{8–12} Because OCT uses broadband light, 2-D and 3-D cross-sectional images of subsurface tissue can be reconstructed at a high resolution for various biological and medical applications.\textsuperscript{13,14} However, OCT-based early cancer detection is limited by the low contrast levels in biological tissues, particularly between normal and neoplastic tissues. Various approaches, including Doppler OCT,\textsuperscript{15–19} polarization-sensitive OCT,\textsuperscript{20–23} chemico-physical enhancers (e.g., absorbing dyes),\textsuperscript{24,25} and contrast agents\textsuperscript{26–29} have been explored to overcome this fundamental limitation of OCT. Little progress has been reported using contrast agents to enhance OCT images \textit{in vivo}.\textsuperscript{27,30} The use of inorganic nanoparticles such as nanospheres,\textsuperscript{31–34} nanocages,\textsuperscript{35,36} nanoshells,\textsuperscript{37,38} and nanorods\textsuperscript{39,40} to overcome OCT limitations, in particular using surface plasmon resonance (SPR), has been investigated, but limited \textit{in vivo} successes have been achieved compared with phantom studies and \textit{in vitro} tests. Some of the limitations for the use of nanoparticles as OCT contrast agents include toxicity [e.g., \textit{in vivo} toxicity of nanorods due to the indispensable use of cetyltrimethylammonium bromide (CTAB) during synthesis]\textsuperscript{41} and poor \textit{in vivo} delivery and distribution.\textsuperscript{42}

Gold nanoparticles (Au NPs) are promising \textit{in vivo} OCT contrast agents because they are biocompatible, easy to synthesize, and functional with additional modalities. For example, Au NPs have been used in human rheumatoid arthritis treatment and microscopy at both low- and high-resolution levels as specific markers for a variety of macromolecules (e.g., polysaccharides, glycoproteins, proteins, lectins, and antibodies).\textsuperscript{43,44} In addition, Au NPs are attractive as OCT contrast agents because the optical resonance properties of Au NPs can be controlled over a broad range by tailoring their sizes and shapes.\textsuperscript{4,45–48} A crossover from a low (absorption dominant) to a high (scattering dominant) albedo of spherical Au NPs at an 80-nm diameter, which can be quantified by OCT, has been theoretically and experimentally proven.\textsuperscript{49}

The efficient delivery of Au NPs is crucial to obtain a sufficient signal and a sensitivity-determining step. A topical administration of Au NPs is preferable because OCT can only image to an approximate depth of 2 mm in human soft tissues.\textsuperscript{50} A topical delivery can avoid adverse systemic effects while delivering a large quantity of Au NPs to the epithelial layers where many types of oral, skin, and gastrointestinal cancers develop. However, a supporting vasculature is poorly developed or not available.\textsuperscript{51} The penetration of Au NPs to subsurface epithelial layers is greatly hampered by various biological barriers such as the stratum corneum.
(SC), the 10- to 15-μm-thick outermost nonliving layer [Fig. 1(a)]. In this study, we hypothesized that this obstacle to Au NP delivery for OCT imaging could be overcome by generating micropassages through the SC using microneedles (MNs), followed by the enhanced distribution of Au NPs in underlying epithelial layers through increased movements by ultrasonic forces (i.e., multimodal delivery) [Figs. 1(b) and 1(c)].

The purpose of this study was to improve the penetration and distribution of Au NPs using MNs and US and thereby enhance contrast in in vivo OCT images of oral dysplasia in a hamster model. Spherical Au NPs were prepared as previously reported and further conjugated with antiepidermal growth factor receptor (EGFR) monoclonal antibodies and polyethylene glycol (PEG), also as previously reported. Au NPs were applied topically with and without 300-μm-long MNs and US, in vivo to the hamster cheek pouch model, and the changes in OCT signals with the same focal point were quantified by intensity profiles.

2 Materials and Methods

2.1 Au NP Synthesis

Au NPs (71 nm in diameter) were prepared by the Frens method with modifications. A 71-nm-diameter Au NP (highly scattering dominant) was used as the OCT contrast agent because OCT detects backscattering signals from a sample. All glassware was cleaned in aqua regia (three parts HCl, one part HNO₃), rinsed with deionized water, and then dried prior to use. Solutions of HAuCl₄ · 3H₂O (0.01% w/w, solution A) and Na₃-citrate (1% w/w, solution B) were prepared, then 100 mL of solution A was heated to its boiling point and 0.6 mL of solution B was added with vigorous stirring. After 25 sec, the boiling solution turned dark blue followed by violet after 2 min of stirring. The solution was refluxed for an additional 15 min and allowed to cool down to room temperature with continuous stirring. The final concentration of 71-nm-diameter Au NPs was 2.19 × 10¹⁰ particles/mL. Particle sizes were determined using a Carl Zeiss Ultra 55 scanning electron microscope (SEM), Phillips CM20 transmission electron microscope (TEM), and Melvern Zetasizer dynamic light scattering (DLS) particle analyzer. The Au NP solution was kept at 4 °C.

2.2 Antibody Conjugation on Au NPs

Solutions of Au NPs were washed three times with deionized (DI) water using centrifugation at 8000 rpm (9230 g) for 30 min. Monoclonal antibodies (Clone 29.1.1, Sigma, St. Louis, MO) binding to EGFR, which is over-expressed in oral cancer, were conjugated on the Au NP surfaces using protocols published elsewhere with modification. Briefly, 1 mL of washed Au NP solution was diluted with 125 μL of 20-mM HEPES buffer (Fisher Scientific, Fair Lawn, NJ) and mixed with 30 μL of an anti-EGFR antibody-containing solution (1 mg/mL) in 70 μL of 20-mM HEPES. The pH of the solution was kept at 7.0 ± 0.2 and stirred for 20 min. The final concentration of the antibody-conjugated Au NP solution was 1.78 × 10¹⁰ particles/mL, and the final volume was 1.225 mL. Then 200 μL of this solution was tested for a color change in the presence of 10 μL of 10% NaCl, since it is well known that NaCl induces Au NP aggregation and results in a change to the solution’s original color. Au NPs that are completely coated with antibodies do not coagulate, so a color change to dark blue indicates the Au NP surfaces are not saturated with antibodies. Finally, 200 μL of a 1% (w/v) PEG (MW=4000) solution (Fluka, Switzerland) in DI water was added, and the mixed solution was incubated for 10 min at room temperature while stirred. The solution was centrifuged at 8000 rpm (9230 g) for 10 min at room temperature, and the pellet was resuspended in 0.5 mL of phosphate buffered saline (Fisher Scientific, Fair Lawn, NJ).
2.3 In vivo Oral Cancer Model and Au NPs Delivery

For in vivo imaging, the standard hamster cheek pouch model was used, whereby the cheek pouch on one side of each golden Syrian hamster (Mesocricetus auratus) was treated topically with 0.5% (v/v) 9, 10-dimethyl-1,2-benzanthracene (DMBA, Sigma, St. Louis, MO) three times per week for five months to induce cancer. The contra-lateral side was used as control, with only mineral oil applied topically to the cheek pouch surface. The anesthetized hamster’s cheek pouch was attached to a microscope stage using a custom-built ring-shaped clamp that was rigidly fastened to the stage surface. CR3 roller microneedles (MTS dermaroller with miniscule holes of 70-μm diameter and 300-μm depth; Clinical Resolution Laboratory, Inc., Beverly Hills, CA) were rolled on both the DMBA-untreated and DMBA-treated sides of the hamster cheek pouches three times at three different angles (i.e., 0 deg, 45 deg, and 90 deg). Two hundred μL of the anti-EGFR antibody-conjugated PEGylated Au NP solution (1.78 × 10^10 particles/mL) was applied to the hamster’s cheek pouch for 10 min by dropping it directly into the 1-cm-diameter aperture of the ring-shaped clamp. After the Au NP topical administration, 0.3 W/cm^2 of 1-MHz ultrasonic force was applied to the cheek pouch using the Dynatron 125 ultrasonicator (Dynatronics Corporation, Salt Lake City, UT) for 1 min. The typical ultrasonic intensity range for diagnostic applications is 0.0001 to 0.5 W/cm^2, and the 1-MHz frequency has been reported to be sufficient to facilitate NP dispersion into tissue by changing the diffusion coefficient (by factors of 2.6 to 15) of the tissue rather than partitioning (factors of 0.7 to 1.6).^{54,58}

2.4 Setup of Spectral-Domain Optical Coherence Tomography

Spectral-domain optical coherence tomography (SD-OCT) is a high-speed, high-resolution, minimally invasive, cross-sectional imaging technique based on a Michelson interferometer with four arms (Fig. 2). The source arm has a broadband light source with a 1310-nm center wavelength and a 90-nm FWHM. The detector arm has a spectrometer with a 0.13-nm spectral accuracy and a 7.7-kHz frame rate. An immobilized mirror is placed at the reference arm. A two-axis scanner with two galvanometers is located at the sample arm. Reflected light from the immobilized mirror and each scattering particle in a sample make a signal in the spectral domain that is detected by the spectrometer. When the path difference between the reference and sample arms is defined by 2y and the refractive index of the sample by n, the scattering event by each particle at a different depth y is encoded in the frequency 2ny of the cosine function, and the signal is a sum of the cosine functions with different y’s, where the amplitude of each cosine is proportional to the scattering amplitude. An inverse Fourier transformation of the signal in the spectral domain gives a complex signal in the y domain. The powers of peaks in the y domain represent the scattering amplitudes and are converted to grayscale to make an array of images along the y direction. While the two-axis scanner scans the xc plane, the array is acquired continuously to make a volume image. The current setup resolution is 8 μm (in air) depthwise and 13 μm laterally. Each pixel is 5 μm × 5 μm in tissue with a refractive index of 1.33. All the SD-OCT images were taken with the same focal point.

2.5 Histology of Excised Tissues

The excised hamster cheek pouch tissues were fixed in Karnovsky 2% paraformaldehyde and 3% glutaraldehyde in 0.1-M cacodylate buffer overnight and then postfix in 1% osmium tetroxide for 1 hr. The tissues were dehydrated in a graded series of ethanol solutions (30%, 50%, 70%, 90%, and 100%), then embedded in PolyBed® 812 (Polyscience, Inc., Warrington, PA) and polymerized at 60 °C for 2 days. Half-μm semi-thin sections were prepared using an ultra microtome and then stained with Richardson’s stain for visualization under an upright microscope (BH-2, Olympus, Tokyo, Japan) equipped with an Olympus Microfire digital camera. To confirm the lack of inflammatory response, MNs, topical delivery of Au NPs, and
US were applied to the cheek pouches of DMBA-untreated hamsters. The cheek pouches were stained with hematoxylin and eosin (H&E) 40 days after the multimodal delivery of Au NPs.

3 Results and Discussion

The results shown in Fig. 3 clearly demonstrate that (1) Au NPs were able to penetrate deep into the tissue only when micropassages were provided by MNs (i.e., Au NP alone versus MN-Au NP and Au NP-US versus MN-Au NP-US for both DMBA-treated and DMBA-untreated tissues in Fig. 3); and (2) US-enhanced Au NP distribution throughout the tissues (i.e., Au NP alone versus Au NP-US and MN-Au NP versus MN-Au NP-US for both DMBA-treated and DMBA-untreated tissues in Fig. 3). In the SD-OCT images, the tissue layers were clearly visible, and the difference between epithelial micromorphologies of the carcinogen-treated and untreated tissues was clearly visible [Figs. 3(a) and 3(b)]. Scion Image software (Scion Corp., Frederick, MA) was used to obtain mean intensities for each SD-OCT image with 894 pixels (length) by 237 pixels (width). The mean intensity of normal tissue was increased by multimodal delivery of Au NPs from 46.64 to 69.25 (149% increased), and the mean intensity of DMBA-treated tissue was increased from 34.59 to 61.08 (177%), as shown in Figs. 3(a) and 3(b), respectively. The images shown in Figs. 3(a) and 3(b) were from the left and right cheek pouches of the same animal. The right cheek pouch was treated with DMBA for five months. In this standard model for oral carcinogenesis, neoplastic lesions will typically begin to develop after four to six weeks in the treated cheek pouch while the untreated cheek pouch is used as the control. However, this animal received the carcinogen for five months. While we had assumed that the untreated cheek pouch could still be used as a healthy control, our OCT images showed the unexpected development of dysplasia in the untreated cheek pouch (dotted box in Fig. 3), which was confirmed by subsequent histopathology.

OCT intensity profiles along with imaging depth were assessed to identify distinctive optical properties of the abnormal tissue (Fig. 4). Five SD-OCT intensity profiles, averaged over 5 pixels laterally, were taken from the SD-OCT image of dysplasia tissue shown in Fig. 4(a). Scion Image software was used for data processing. The actual SD-OCT image depth was calculated by converting the axial pixel number to actual scale by multiplying 5 μm for each pixel in the tissue by a refractive index of 1.33. Several quantitative OCT markers of dysplasia were also identified: (1) considerably greater light scattering in the SC and upper epithelial layers [red square in Fig. 4(b)]; (2) reduced optical signals in subsurface epithelial layers [green square in Fig. 4(b)]; and (3) relatively low OCT signals in underlying connective tissue layers [blue square in Fig. 4(b)].

SD-OCT-based diagnoses (shown in Figs. 3 and 4) were verified by histological staining (Richardson’s stain) of the excised tissues. Figure 5 shows that Au NPs were dispersed along passages generated by the 300-μm-long MNs. Au NPs were dispersed homogeneously into the connective tissue [Fig. 5(a)] but not into the hamster cheek pouch muscle bundles [Fig. 5(b)]. This may be due to the presence of a muscle outer layer that prevented penetration of the Au NPs. The greater presence of Au NPs in connective tissue rather than muscle bundles, as indicated by strong backscattering signals (Figs. 3 and 4), was confirmed by histology. Interestingly, Au NPs were able to penetrate deeper through the epithelial layers in the premalignant region [dotted area in Fig. 6(a)] than in the normal region [Fig. 6(b)]. This finding explains the increased signals in the SC and upper epithelial layers in the early dysplasia region [red square in Fig. 4(b)]. Arrows in Fig. 6(a) indicate the disrupted epithelial layers of an early-stage dysplasia, thus demonstrating that mild dysplasia was differentiated from normal tissue by SD-OCT with enhanced Au NP distribution.

The duration of Au NP-induced OCT signal enhancement was also investigated (Fig. 7) by imaging the cheek pouches of DMBA-untreated hamsters over a period of 40 days after the
initial administration using MNs and US. No noticeable changes were observed for two days. OCT signals in epithelial and connective tissue areas were found to be significantly diluted after a week, indicating that the Au NPs were cleared from the lower tissue. However, enhanced signals from the SC persisted over the entire 40 days. This may be due to poor (or no) active transport in the SC area. Thus, it was speculated that Au NPs were efficiently delivered by multimodal methods to both the epithelial and connective layers, where relatively active clearance of Au NPs is available. This could be confirmed by quantifying the concentration of Au NPs in the tissue. Importantly, no inflammation or other adverse effects were observed in the hamster cheek pouch over a period of 40 days, indicating the potential for an acceptable level of biocompatibility of the Au NPs (Fig. 8). PEGylated Au NPs with an antibody coating were used to prevent agglomeration of Au NPs while achieving EGFR-mediated specific binding to cancerous cells. Au NP agglomeration in vivo was effectively prevented, but no selective antibody-mediated binding of the Au NPs was observed in these studies (i.e., the low optical signals in the dysplasia shown in Figs. 3 and 4, and no selective accumulation of Au NPs in the dysplasia membrane shown in Fig. 6). This may be explained by the predominant enhanced permeation and retention (EPR) effects on relatively large particles in tissue. Due to their size (i.e., 71 nm), Au NPs may have been immobilized in the tissue with a low EGFR-encountering frequency. This unanswered question should be investigated in a subsequent study with a quantitative model and experimental data using nonspecific antibody-conjugated Au NPs. The antibody seemed to at least serve as a PEGylation anchor via its strong affinity to Au surfaces and efficient absorption of PEG on antibody layers.

4 Conclusion

In conclusion, our multimodal delivery that employed MNs and US enabled the efficient transport of Au NPs by overcoming the SC and epithelial barriers and resulted in an approximately 150% increased OCT contrast level in the standard model for oral carcinogenesis. Early neoplasia was clearly identified by effectively delivered and distributed Au NPs, which also permitted meaningful quantitative signal analysis. The distribution of Au NPs in subepithelial layers through MN-assisted penetration and US-facilitated distribution was confirmed by histology. Diminishing OCT signals in sub-SC layers over time (up to 40 days) imply that Au NPs were able to reach the area within the tissues where active and efficient clearance is available. This study demonstrates an effective approach to overcoming poor transport of OCT contrast-enhancing nanoparticles and may provide a new paradigm for enhancing in vivo OCT images for the early diagnosis of cancer. Further study on the molecular properties of Au NPs (e.g., Au NP size, shape, and surface functionalities) and on systemic tuning of multimodal delivery methods (e.g., MN geometry, level and duration of ultrasonic forces, and various combinations of the two modalities) are underway to further elucidate and optimize this new approach.

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Fig. 1.
Stratum corneum and underlying epithelial layers as biological barrier to Au NP transport. (b) Penetration of Au NPs into subsurface epithelial layers through a passage generated by a microneedle. (c) Enhanced Au NP distribution in tissue by ultrasonic force.
Fig. 2.
Schematic of a fiber-based SD-OCT in which the lowcoherence light has a 1310-nm center wavelength and 90-nm FWHM. A 130-nm-wide spectrum was sampled by a 1 × 1024 InGaAs detector array at 7.7 kHz. Imaging depth and depth resolution were 3.4 mm and 8 μm in air, respectively. A two-axis scanner with two galvanometers was used. CM: collimator; DAQ: data acquisition system; DG: diffraction grating; FL: focusing lens; LCL: low-coherence light; LSC: line scan camera.
Fig. 3.  
*In vivo* SD-OCT images of (a) normal (DMBA-untreated), and (b) DMBA-treated sides of hamster cheek pouches. MN = microneedle treated; Au NPs = gold nanoparticles administered; US = ultrasound applied.
Fig. 4.
(a) Enlarged OCT image of dysplastic area. (b) Depth-resolved OCT signal profile in dysplastic and normal regions. Scale bar: 100 μm.
Fig. 5. Au NPs in a hamster cheek pouch tissue after MN and US administration. Arrows indicate a MN passage in the tissue.
Fig. 6.
Distribution of Au NPs in SC and upper epithelial layers of (a) an early dysplasia, and (b) a normal region.
Fig. 7.
Kinetics of in vivo OCT signal profiles in a normal hamster cheek pouch for 40 days. Scale bar: 100 μm.
Fig. 8.
H&E stained cheek pouch of a DMBA-untreated hamster 40 days after MNs, Au NPs, and US were applied.