Surface-Chemistry Effect on Cellular Response of Luminescent Plasmonic Silver Nanoparticles

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Supporting Information

ABSTRACT: Cellular response of inorganic nanoparticles (NPs) is strongly dependent on their surface chemistry. By taking advantage of robust single-particle fluorescence and giant Raman enhancements of unique polycrystalline silver NPs (AgNPs), we quantitatively investigated effects of two well-known surface chemistries, passive PEGylation and active c-RGD peptide conjugation, on in vitro behaviors of AgNPs at high temporal and spatial resolution as well as chemical level using fluorescence and Raman microscopy. The results show that specific c-RGD peptide–αvβ3 integrin interactions not only induced endosome formation more rapidly, enhanced constrained diffusion, but also minimized nonspecific chemical interactions between the NPs and intracellular biomolecules than passive PEGylation chemistry; as a result, surface enhanced Raman scattering (SERS) signals of c-RGD peptides were well resolved inside endosomes in the live cells, while Raman signals of PEGylated AgNPs remained unresolvable due to interference of surrounding biomolecules, opening up an opportunity to investigate specific ligand–receptor interactions in real time at the chemical level.

INTRODUCTION

Quantitative understanding of surface-chemistry effects on in vitro behaviors of inorganic nanoparticles (NPs) is of fundamental importance to their future success in preclinical research and clinical practices.1–12 Two surface chemistries, PEGylation and conjugation of active targeting ligands such as cyclic Arg-Gly-Asp (c-RGD) peptides, have been widely applied to enhance the delivery of NPs into tumor sites.8,13–15 With the assistance of PEG molecules, NPs can avoid macrophage uptake and passively accumulate in tumors at high efficiency through well-known enhanced permeability and retention (EPR) effect.16–18 For instance, PEGylated gold NPs with a core size of ~2.3 nm exhibited high tumor targeting efficiencies of 8.3 ± 0.9%ID/g at 12 h post injection (p.i.).18 Gold nanocages coated with PEG surface ligand also enabled high blood retention and an efficient tumor uptake of 7.9 ± 1.1%ID/g at 24 h p.i.17 Alternatively, conjugation of active targeting ligands that exhibit high affinities to cancer receptors is also a widely used strategy to increase tumor-targeting efficiencies of the NPs.5,19 For example, c-RGD coated iron oxide NPs targeted integrin expressing tumor vasculature after injection, and a significant decrease of tumor magnetic resonance (MR) signal intensity (42 ± 5%) was observed.20 While both surface strategies can enhance the tumor delivery of NPs at the in vivo level, how these two different surface chemistries quantitatively influence their follow-up cellular uptake, intracellular dynamics, and local chemical interactions inside endosomes still have not been fully understood and demand substantial investigations. To address these questions, NPs that not only exhibit robust single-particle fluorescence but also can chemically report local interactions are highly desired.

Luminescent plasmonic nanoparticles (LPNPs) are a class of nanofluorophores which can give bright and robust single-particle fluorescence while exhibiting plasmon properties such as strong surface plasmon absorption and large surface-enhanced Raman scattering (SERS) enhancements.21–23 The synergy of these properties in one NP allows strengths of different optical microscopic imaging techniques to be integrated for better understanding of nano-bio interactions at the in vitro level. One strategy for creating such LPNPs is to coat a fluorescent quantum dot (QD) with an ultrathin gold nanoshell through precise control of the distance between QD and NPs.
and the shell with elegant surface chemistries. Alternatively, by tuning grain size distributions of metal NPs, we were able to create 20 nm highly polycrystalline silver or gold NPs with strong single-particle fluorescence and giant Raman enhancements. While simple cellular imaging of cancer receptors labeled with LPNPs has been demonstrated, how to further apply them to advance our fundamental understanding of nanobio interactions has not yet been demonstrated. In this communication, we applied luminescent plasmonic silver NPs (LPAgNPs) as contrast agents to quantitatively investigate the effects of two widely used surface chemistries in cancer targeting, passive PEGylation, and active c-RGD conjugation, on cellular uptake, intracellular dynamics, and local chemical interactions of the NPs inside the endosomes. Our studies showed that the conjugation of c-RGD peptides (Figure S1) to the NPs resulted in 20 times more rapid cellular uptake and also enhanced constrained motion of the NPs about 13% higher than that of PEGylation. In addition, specific binding between c-RGD and \( \alpha_v \beta_3 \) integrin minimized interference of intracellular biomolecules on Raman signals of LPAgNPs and allowed c-RGD vibrations to be well dissolved inside endosomes in the live cells, offering an exciting opportunity to investigate ligand–receptor interactions in real time at the chemical level.

**RESULTS AND DISCUSSION**

The LPAgNPs with a mean size of \(~20 \text{ nm}\) were selected for the follow-up bioconjugation and *in vitro* cellular studies. To conjugate c-RGD peptides or PEG molecules onto the particle surface, the LPAgNPs were incubated with mercaptobenzoic acid (MBA) and c-RGD peptides or 1 kDa PEG molecules. MBA ligand was introduced as the secondary ligand because its characteristic vibrations at 1071 cm\(^{-1}\) and 1575 cm\(^{-1}\) can serve as an internal standard during SERS studies. After the bioconjugation, the obtained NPs were further characterized with transmission electron microscopy (TEM) and dynamic light scattering (DLS). DLS studies showed that the hydrodynamic diameters (HDs) of the NPs coated with either c-RGD peptides or PEG molecules were 28.1 ± 6.1 nm (Figure 1a) and 33.7 ± 6.9 nm (Figure 1b) in phosphate buffer saline (PBS), respectively, indicating that NPs remained monodispersed in the physiological environment. HD layer of PEG-LPAgNPs (\(~7 \text{ nm}\)) was slightly larger than that of c-RGD conjugated ones (\(~4 \text{ nm}\)), indicating that PEG exhibited an

Figure 1. Characterization of LPAgNPs. (a) c-RGD-LPAgNPs with a core size of 20.9 ± 4.6 nm, and a hydrodynamic diameter (HD) of 28.1 ± 6.1 nm. (b) PEG-LPAgNPs with a core size of 20.5 ± 4.2 nm, and a HD of 33.7 ± 6.9 nm. The scale bar of the TEM images is 50 nm. (c) Absorption spectra of c-RGD-LPAgNPs and PEG-LPAgNPs. (d) Emission spectra of c-RGD-LPAgNPs and PEG-LPAgNPs. (e) Raman spectra of c-RGD-LPAgNPs. (f) Raman spectra of PEG-LPAgNPs.
extended conformation once coated on the NP surface. In addition, zeta potential measurements on these NPs showed that c-RGD-LPAgNPs and PEG-LPAgNPs were negatively charged in PBS buffer with zeta potentials of $-30$ mV and $-32$ mV, respectively. The obtained AgNPs retained their strong surface plasmons at 400 nm and exhibited identical absorption profiles, further suggesting that conjugation of the different ligands on the NP surface induced little change on their absorption (Figure 1c). Emission spectra of both PEG-LPAgNPs and c-RGD-LPAgNPs were composed of broad fluorescence background superimposed with sharp Raman emissions. Since PEG and c-RGD exhibit different Raman vibrational spectra, the overall emission spectra of AgNPs conjugated with different ligands exhibit different profiles (Figure 1d). The observation of Raman spectra from c-RGD-LPAgNPs and PEG-LPAgNPs also indicated that surface ligands were successfully conjugated on the NP surface. As shown in Figure 1e,f, Raman spectra of c-RGD-LPAgNPs showed the characteristic Raman vibrations of c-RGD ligands on PEG-LPAgNPs such as the $\beta$-sheet (1631 cm$^{-1}$), II (1518 cm$^{-1}$), III (1265 cm$^{-1}$), and phenyl ring (1179, 1212 cm$^{-1}$), of c-RGD were readily observed, consistent with the previous report. Characteristic vibrations of PEG ligands on PEG-LPAgNPs such as the $\mathrm{CH}_2$ rocking vibration (936 cm$^{-1}$), $\mathrm{C}=$O–C asymmetric stretching (1129 cm$^{-1}$), the coupled vibration of $\mathrm{C}–\mathrm{C}$ and $\mathrm{C}–\mathrm{O}$ stretching (1132 cm$^{-1}$) and $\mathrm{CH}_2–\mathrm{CH}_2$ antisymmetric bending mode (1462 cm$^{-1}$) were also observed from PEG-LPAgNPs. In addition, Raman bands of MBA on the LPAgNPs were observed at 1071, 1352, and 1575 cm$^{-1}$, which were exactly consistent with the previous report on C–C stretching ring-breathing modes (1071 and 1575 cm$^{-1}$) and COO stretching (1352 cm$^{-1}$), respectively. 

LPAgNPs with different surface ligands exhibited distinct binding affinities to glioblastoma U87MG cancer cells with high expression level of $\alpha_\beta_3$ integrin receptors on the membrane. The fixed U87MG cancer cells were incubated with c-RGD-LPAgNPs and PEG-LPAgNPs of the same concentration (10 nM) in PBS respectively. After 45 min incubation, the excess amount of NPs was washed away with PBS and the cells were imaged under a fluorescence microscope with a 100× objective in total internal reflection geometry. As shown in Figure 2a,b, only a very small amount of PEG-LPAgNPs was found to nonspecifically bind to the U87MG cells because of the antifouling nature of PEG molecules. On the other hand, a large amount of c-RGD-LPAgNPs was labeled on the cell membrane (Figure 2c,d). Because of a strong surface plasmon of c-RGD-LPAgNPs and their high-density labeling of U87MG cancer cells, the cells became yellowish and could even be readily observed with the naked eye (Figure S2). To further confirm that the observed high labeling efficiency of c-RGD-LPAgNPs was due to specific ligand–receptor interactions, we also conducted blocking and c-RAD peptide control studies. We first blocked the binding of the integrin receptors of U87MG cells with 1 mM c-RGD peptide and then incubated U87MG cells with c-RGD-LPAgNPs with the same procedures. No binding of the AgNPs with U87MG cells was observed (Figure 3a,b). In addition, we also incubated U87MG cells with LPAgNP conjugated c-RAD, a widely used negative control peptide for c-RGD, and observed very little binding of the AgNPs to the cells (Figure 3c,d). With all these control studies, we could indeed attribute the high labeling efficiency of c-RGD-LPAgNPs to the high binding affinity of c-RGD peptides to $\alpha_\beta_3$ integrin receptors. These results also indicated successful conjugation of c-RGD to LPAgNPs, and the conjugated c-RGD peptides retained their high binding affinity and specificity to cancer receptors on the membrane. 

Once LPAgNPs (1 nM in MEM) were incubated with live U87MG cancer cells at 37°C, the NPs were internalized by the cells, where the NPs were encapsulated in endosomes regardless of which types of ligands were on the surface. To gain a quantitative understanding of the differences in endocytosis kinetics between two types of LPAgNPs with
different surface coatings, we used fluorescence microscopy to quantify the average number of endosomes at different incubation time points. The number of endosomes incubated with c-RGD-LPAgNPs reached a platform with 49 ± 9 endosomes observed at the first 3 h (Figure 4a and Figure S3b), while only very few endosomes (2–3) containing PEG-LPAgNPs were found inside cells during the same time period (Figure 4b and Figure S3a), suggesting the specific ligand−
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α appear in the same spectral region),
derived from amino acid residues and proteins are broad and
be due to the signi-
recovered (Figure S5). Such broadening in Raman spectra can
PEG, c-RGD, and MBA ligands on the particle surface were
centrifugation puri-
in the presence of MEM cell culture media (Figure 5a,b) or
in PBS containing 10% FBS (v/v) (Figure S4). However, once the
in the media (the neighboring bands
RGD-LPAgNPs and PEG-LPAgNPs in PBS solution (Figure
−1 and 1515 cm
while
observed from c-RGD-LPAgNPs indicated that the speci-
LPAgNPs exhibited directed motion, while ~10% and ~23% showed
Brownian diffusion and constrained diffusion, respectively (Figure 4c).
On the other hand, ~67% of endosomes encapsulated PEG-
characteristic Raman vibrations of c-RGD-LPAgNPs and resulted in recovery
interestingly, speci-
formaldehyde for 10 min. Then the
3 integrin receptors potentially can minimize
interference from local intracellular biological molecules.

### EXPERIMENTAL PROCEDURES

#### Synthesis of Luminescent Plasmonic Silver Nanoparticles (LPAgNPs)
LPAgNPs were synthesized using the solid-state thermal reduction method we reported before.23 200 mg of glycine and 15 mg of silver nitrate were codissolved in 1.5 mL DI water. After the water evaporated, the mixture was then reduced at 453 K for 2 min in the solid state. When the color of the mixture became blackish, the reaction was stopped and the product was dissolved in 1 mL of DI water. The final solution was first centrifuged at 2000 and 4000 g for 2 min respectively to remove the large aggregates. LPAgNPs of ~20 nm were obtained by collecting the pellets after centrifuging the supernatant at 7000 g for 3 min. The sample was further purified by centrifugation at 7000 g several times to remove the free ions and excess glycine ligands. The pellets were collected and redissolved in 10 mM sodium borate buffer for further study.

#### Bioconjugation of LPAgNPs
Mercaptobenzoic acid (MBA), as a ligand to serve as the internal standard of SERS studies, was dissolved in methanol solution (10 mM) and diluted to 1 mM using 10 mM sodium borate buffer. The conjugation of c-RGD peptides, cyclo (Arg-Ala-Asp-D-Phe-Cys) (c-RAD) peptides, or PEG molecules to LPAgNPs was conducted by incubating the LPAgNPs with MBA and c-RGD (c-RAD, PEG) at a molar ratio of 1:5 × 10^{-15}:5 × 10^{4} in 10 mM sodium borate buffer for 2 h with gentle shaking. The excess ligands and free ions were removed by passing the sample through a Sephadex-20 column with PBS as elution phase. The sample was then centrifuged at 7000 g for 3 min and the pellets were redissolved in phosphate buffer saline (PBS) for further study.

#### Fixed Cell Labeling and Imaging
For cell labeling, U87MG cells were first fixed by 3.7% formaldehyde for 10 min. Then the fixative was removed and the cells were blocked with 1 mL 2 weight (wt) % BSA for 30 min. The fixed cells were incubated with 10 nM c-RGD-LPAgNPs, c-RAD-LPAgNPs, or PEG-LPAgNPs in 500 μL PBS buffer solution at room temperature for 45 min, respectively. The cells were then washed thoroughly with PBS buffer to remove the excess amount of NPs and used for imaging studies.

#### Blocking Study
U87MG cells were first fixed by 3.7% formaldehyde for 10 min. Then the fixative was removed and the cells were blocked with 1 mL 2 wt % BSA for 30 min. The fixed cells were incubated with 1 mM c-RGD in 500 μL PBS buffer solution at room temperature. After 45 min, the c-RGD was removed and the cells were washed thoroughly with PBS buffer for 3 times. Then the cells were incubated with 10 nM c-RGD-LPAgNPs for another 45 min before being used for cell imaging studies.

#### Cellular Endocytosis Studies
The live cells were incubated with 1 nM c-RGD-LPAgNPs in MEM for 0.5, 1,
1, 5, 2, 2.5, and 3 h or with 1 nM PEG-LPAgNPs in MEM for 3, 6, 12, and 24 h at 37 °C, 5% CO₂. The cells at different time points were washed thoroughly with PBS buffer and used for cell imaging studies. Before the fluorescence cell images were taken, the cells were photographed for 15 s. For each time point, the bright endosomes in each cell were localized and counted by Insight software developed by Dr. Bo Huang (Department of Pharmaceutical Chemistry and Department of Biochemistry and Biophysics, University of California, San Francisco). The size of the bright dots was chosen as larger than 50 pixels. Ten cells were analyzed at each time point and presented as mean ± SD (n = 10).

**Cellular Dynamic Studies.** The live cells incubated with c-RGD-LPAgNPs for 3 h and the live cells incubated with PEG-LPAgNPs for 24 h were used for the cellular dynamic studies, respectively. After 15 s photobleaching, fluorescence cell images were taken every 500 ms for 50 s. The Insight software was used for the localization and tracking of the bright dots (endosomes). The bright dot size was chosen as larger than 50 pixels. For the tracking study, the minimum trace length was 20 frames. The motions of the NPs were classified into three different modes according to the α (slope) value of the trajectories: constrained diffusion (α < 0.9), Brownian diffusion (0.9 ≤ α ≤ 1.1), and directed motion (α > 1.1). [31]

**Cellular SERS Studies.** U87MG cancer cells were incubated with c-RGD-LPAgNPs or c-RAD-LPAgNPs for 3 h, or with PEG-LPAgNPs for 24 h. The cells were washed thoroughly with PBS for 3 times after incubation to remove the excess amount of LPAgNPs. The Raman spectra were then collected with an Acton SP2300 (Princeton Instruments) under 532 nm excitation.

**REFERENCES**


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