Dealloyed Intra-Nanogap Particles with Highly Robust, Quantifiable Surface-Enhanced Raman Scattering Signals for Biosensing and Bioimaging Applications

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Supporting Information

ABSTRACT: Uniformly controlling a large number of metal nanostructures with a plasmonically enhanced signal to generate quantitative optical signals and the widespread use of these structures for surface-enhanced Raman scattering (SERS)-based biosensing and bioimaging applications are of paramount importance but are extremely challenging. Here, we report a highly controllable, facile selective-interdiffusive dealloying chemistry for synthesizing the dealloyed intra-nanogap particles (DIPs) with a ~2 nm intragap in a high yield (~95%) without the need for an interlayer. The SERS signals from DIPs are highly quantitative and polarization-independent with polarized laser sources. Remarkably, all the analyzed particles displayed the SERS enhancement factors (EFs) of \( \geq 1.1 \times 10^8 \) with a very narrow distribution of EFs. Finally, we show that DIPs can be used as ultrasensitive SERS-based DNA detection probes for detecting 10 aM to 1 pM target concentrations and highly robust, quantitative real-time cell imaging probes for long-term imaging with low laser power and short exposure time.

INTRODUCTION

Plasmonic nanostructures, particularly with a plasmonically coupled and enhanced nanogap, are of paramount importance and interest due to their strong, tunable optical properties and enormous potential in various applications including catalysis, sensing, and imaging.1−4 Because the optical properties of plasmonic nanostructures are highly dependent on various nanostructural features, highly precise synthesis of plasmonic nanostructures in a high yield is needed to obtain controllable, quantifiable plasmonic signals from these nanostructures. Among many plasmonically enhanced optical signals, surface−enhanced Raman scattering (SERS) has received lots of attention because of its ultrahigh sensitivity, which can be extended to single-molecule-level sensitivity and multiplexing potential.5 In this regard, SERS-based techniques are being exploited widely in plasmonics and analytical applications, including for biosensing and the detection of chemical warfare agents.6,7 In plasmonics, it is now well-known that the excitation of localized surface plasmon resonances in nanoparticles (NPs) of noble metals (in particular, those of Au and Ag) can significantly enhance and localize the electromagnetic (EM) field at specific locations between NPs, which are called “hot spots”.8−10 There have been numerous attempts to design and produce SERS-active structures by using morphology-controlled noble metal nanostructures, such as nanotips, nanopores, and nanoscale-roughened surfaces.11−13 In particular, with respect to SERS, the EM field can also be enhanced significantly via plasmonic coupling at the gaps or junctions between metallic nanostructures.1 However, the plasmonic coupling in these gaps is largely affected by the interparticle distance, which, in turn, has a marked effect on the SERS signal.15 This can induce uncontrollable irreproducibility in the SERS signal, which is a major barrier for these SERS-active structures to be useful for practical applications. Consequently, a high-precision nanometer scale or sub-nanometer scale gap engineering for a large number of these nanostructures is a key challenge for generating reproducible and reliable SERS signals from these particles.16−17

Among many plasmonic nanogap structures,1 intra-nanogap structures with a nanometer-scale interior gap are highly promising as SERS substrates because of their uniformity and controllability in nanogaps over a large number of particles, resulting in strongly enhanced and tunable SERS signals. To form SERS-active interior nanogap structures, various synthetic strategies have been adopted.18−20 In particular, it was shown that thiolated DNA on AuNPs can facilitate the formation of a 1 nm interior gap during the Au−Au core−shell formation process and generate Au nanobridged nanogap particles (AuNNPs) with a uniform 1 nm intragap in a high yield and strong...
stable, and reproducible SERS signals. Other approaches in generating interior nanogap structures such as silica(SiO$_2$)-interlayered nanogap structures with Au core and Au shell (Au nanomatryoshka) and polymer or small molecule-interlayered plasmonic nanogap structures have been adopted to synthesize various intra-nanogap structures. All these structures,
however, still need to address many drawbacks and challenges such as complicated synthetic complexity, time and cost, reaction conditions including temperature, nanometer-level structural precision, limitation in large-scale production, particle stability, versatility in structure and composition, and lack of practical applications. It should be noted that the need for an interlayer to form intra-nanogaps inside particles and laser polarization-dependent changes in optical signals from intra-nanogap particles largely limit understanding and controlling the plasmonic properties of these nanostructures including electromagnetic field intensity and distribution. Importantly, the use and advantageous properties of these intra-nanogap particles for biomedical applications have not been well explored nor clearly shown.

Here, we present a straightforward, facile and generally applicable synthetic strategy for forming dealloyed intra-nanogap particles (DIPs) with an interlayer-free intra-nanogap in a high yield via the selective-interdiffusive dealloying (SID) process at room temperature, and it is shown that these DIPs can generate strong, quantifiable laser polarization-independent SERS signals (Figure 1A). Importantly, biologically functionalized DIPs can be used for detecting DNA in a highly sensitive and quantitative manner and also for targeting and imaging cancer cells for highly reliable, quantitative, and long-term subcellular bioimaging applications. As the highly controllable dealloying reaction progresses with the DIPs, a nanometer-sized interior gap was formed on Au core surface via selective Ag-etching and interdiffusion of Ag atoms (SID process), resulting in the formation of NPs with a ~2 nm interior gap in high yield (~95%). It should be noted that this SID process can spontaneously happen at room temperature due to the nature of the corrosion process occurring in the presence of etchants or oxidants even at the low temperature. Remarkably, the interior nanogap was uniformly formed with randomly distributed metal residues in a symmetric fashion, leading the polarization-independent SERS signals from the Raman dyes in the interior gaps of these structures, and the SERS signal intensity linearly increases as the particle concentration increases, allowing for highly reliable, quantitative SERS. Remarkably, all the particles (110 particles) displayed SERS enhancement factors of ≥1.1 × 10^6 with a very narrow distribution of EFs. Importantly, Au-gap-Au/Ag DIPs were modified with DNA for DNA sensing applications, and the assay results show that these DNA-modified DIPs generate highly quantitative and sensitive DNA detection results, ranging from aM to pM target concentrations. Finally, these DIPs were functionalized with cell-targeting peptides for cell imaging applications. The results suggest that the peptide-modified DIPs can selectively target integrin αvβ3 expression cells and allow for in situ long-term (>30 min) real-time SERS imaging intracellular regions with low laser power and short exposure time in a highly reliable, quantifiable manner.

■ RESULTS AND DISCUSSION

Selective, Interdiffusive Dealloying-Based Synthesis of Nanoparticles with an Interlayer-Free Interior Nanogap. To synthesize the DIPs with the SID process, we first formed Au/Au−Ag core/ alloy shell nanoparticles (CAS NPs) through a coreduction process involving the simultaneous reduction of HAuCl₄ and AgNO₃ on 4-MPy-modified AuNPs (MPy-AuNPs) (Figure 1B). The average size of CAS NPs, as measured from transmission electron microscopy (TEM) images, was ~87.0 nm (Figure S1), while the atomic composition of the shell region was estimated to be Au₇₇.₃Ag₂₂.₇ (Table S1). Interestingly, the reduced Au and Ag atoms were not evenly distributed throughout the entire shell region (Figure 1C and Figure S2A). An energy-dispersive X-ray spectroscopy (EDX) line scan performed across the centers of the CAS NPs showed that most of the Ag atoms were located near the Au core, with the number of Au atoms decreasing near the Au core (Figure 1D, the blue dotted box). In general, a higher standard reduction potential of the Au precursor results in faster reduction as compared to the case for the Ag precursor (AuCl₄⁻/Au = 0.99 V and Ag⁺/Ag = 0.8 V versus standard hydrogen electrode). The strong affinity between Ag and the pyrrolidine groups of polyvinylpyrrolidone (PVP) help increase the reduction rate of the Ag precursor in the early stage, which results in the accumulation of Ag atoms at a faster rate near the Au core. This result is in good agreement with previously reported results. To examine this phenomenon in more detail, we monitored the changes in the extinction peak of the CAS NPs during the alloy shell formation process (Figure S3). In the early stage of the formation process, the extinction peak exhibited a rapid blue shift, suggesting the formation of Ag layers. As the reaction progressed, the extinction peak underwent a gradual red shift, with the intensity of the peak increasing; this was indicative of the formation of larger CAS NPs.

Next, we introduced ferric nitrate [Fe(NO₃)₃] into the CAS NPs to selectively dissolve the Ag atoms (Figure 1A). It can be seen from Figure 1E that the Ag-etched CAS NPs have a nanogap present between the core and the shell via the dealloying process. The average size of the DIPs (~87.2 nm) was similar to that of the CAS NPs, with the interior nanogap and the shell size of the DIPs being ~2.1 and 20.1 nm, respectively (Figure 1K). The EDX elemental maps of the DIPs confirm that the Ag atoms near the Au core were almost completely etched away (Figure 1F and Figure S2B), and the EDX line scan profile indicates that the number of Ag atoms near the Au core dramatically decreased after the dealloying reaction (Figure 1G, blue dotted box); the proportion of Ag atoms in the shell region was lower as compared to that in the CAS NPs (Table S1). Furthermore, the Au-La curve for the area near the Au core was valley-shaped, when the Ag-La peak disappeared (Figure 1G, blue dotted box), confirming that the interior nanogap is formed by the highly selective removal of Ag atoms near the Au core. In contrast to the case for the DIPs, an interior nanogap was not observed in the case of the gap-less Au/Au core/shell NPs (gap-less AuNPs), whose shell was composed only of Au and not a Au−Ag alloy (Figure 1H). The average size of the gap-less AuNPs was also similar to that of the CAS NPs (Figure S1), with the shell being composed entirely of Au atoms (Figure 1, Figure S2C and Table S1). Further, a valley-like Au-La plot was not observed (Figure 1J, blue dotted box), confirming that the NPs had a nanogap-less structure. When Fe(NO₃)₃ was introduced into the as-synthesized gap-less AuNPs, there was no change in the morphology or the ultraviolet–visible (UV–vis) and SERS spectra of the NPs, suggesting that the ferric nitrate-based dealloying reaction only dissolves Ag (Figure S4). This synthesis strategy led to the fabrication of DIPs in a yield of ~95% (Figure 1L). The δ-spacing of the adjacent lattice fringes of the DIPs was 0.235 nm, and their selected-area electron diffraction (SAED) was ring-like; the former corresponded to the (111) planes of a face-centered cubic structure while the latter was indicative of polycrystallinity (Figure 1M).
To investigate the evolution of the interior nanogap, we monitored the structural changes as the CAS NPs transformed into DIPs through the dealloying reaction while increasing the amount of Fe(NO₃)₃ added (Figure S5). The mechanism proposed to explain the formation of the interior-nanogap structure is shown below (Figure 1 A, see the proposed mechanism in the black dotted box). Once the Fe(NO₃)₃ solution is injected into the CAS NPs, the Ag atoms near the shell surface start getting dissolved by the Fe(NO₃)₃ (the gray line in a schematic of the proposed mechanism), and Fe(NO₃)₃ penetrates the CAS NPs through the Ag-etched sites. As the dealloying reaction progresses, pinhole-like vacancies (the black dotted open circles in the proposed mechanism) di ffuse inward (the blue lines in the proposed mechanism), while metal atoms (mainly Ag atoms) di ffuse outward because of the nanometer-scale Kirkendall effect.²⁷ Because the diffusion of Ag in Au is faster than that of Au in Ag in the case of the Au–Au alloy system, the net flux of the metals (Au and Ag) is mainly dependent on the diffusion rate of Ag, this results in a net flux of vacancies from the shell surface to the Au core.²⁷ Thus, the diffused Ag atoms are continually removed by the Fe(NO₃)₃ (the gray dotted lines in the proposed mechanism). Meanwhile, Au atoms diffuse to a smaller degree and interconnect with each other to minimize their surface energy. As this reaction continues, the accumulated vacancies near the Au core eventually result in the formation of a nanogap, while the interconnected Au atoms form a dense shell. In similarity to the proposed mechanism, the Ag atoms were partially etched and formed an incomplete interior nanogap when a small amount of Fe(NO₃)₃ (5 mM) was used. However, the number of dissolved Ag atoms was higher (10 mM), and an interior nanogap was uniformly formed when a large amount of Fe(NO₃)₃ was used (>15 mM) (Figure S5).

Owing to the morphological differences between the structures with (DIPs) and without (CAS NPs and gap-less AuNPs) a nanogap, the UV–vis spectra of these structures were also different (Figure 2A). Further, with formation of the shell structure, the color of the particle solution changed from pink (MPy-AuNPs) to dark pink (CAS NPs), then to blue-violet (DIPs), and finally to red-violet (gap-less AuNPs), as shown in Figure 2A. Furthermore, the extinction peaks of the NPs with the shell structure were red-shifted compared to those of the shell-less NPs (λ = 528, 549, 554, and 571 nm for the MPy-AuNPs, CAS NPs, DIPs, and gap-less AuNPs, respectively); however, a new extinction shoulder peak was observed at ∼700 nm in the case of NPs with the interior nanogap (DIPs). To elucidate the origin of this plasmonic peak from DIPs and study the optical properties of the nanogap region, we performed a computational simulation based on the Mie theory.²⁸ In our model, metal residues exist and are randomly distributed in the intra-nanogap region, the effective dielectric function of the gap region was calculated using the Smith’s approach that describes metallic behavior of mixtures with low metallic volume fraction (see details in Supporting Information).²⁹ A new extinction peak appears at a
longer wavelength (∼950 nm) when an interior nanogap, filled with water, is formed inside an NP (Figure S6). When the intra-nanogap region is filled with a mixture of metal residues and water, however, the resonance peak was blue-shifted from a longer wavelength region by changing the optical/physical state (e.g., effective dielectric function) of the nanogap. In addition, the resonance peak was continuously red-shifted as metal composition in the nanogap decreased (Figure 2B). This trend is well matched with changes in the UV−vis spectra of DIPs for different degrees of the dealloying reaction (Figure S5). Since the degree of etching for each DIP in solution should be slightly different, the broadened shoulder peak from the experimental result can be explained by the superposition of UV−vis spectra of DIPs with different metal compositions in the nanogap.

Figure 2C shows the calculated near-field EM field distribution of the DIP containing 12.5 mol % of metal residues within the nanogap. Unlike the DIPs with a water-filled intra-nanogap (Figure 2D), the DIPs with a metal residue-filled nanogap generated a highly enhanced EM field, indicating that the metal residues within the interior nanogap play an important role in EM field enhancement.

**SERS Analysis of DIPs in Solution.** Next, we compared the solution-state SERS signals obtained from the as-synthesized NPs with respect to the fingerprint peak at 1097 cm^-1 (Figure 3A). As compared to the SERS signal intensity of the shell-less Au NPs (MPy-AuNPs), those of the CAS NPs, gap-less AuNPs, and DIPs, were ∼8.6, 7.7, and 50 times higher, respectively. In this case, Raman reporter molecules (4-MPy) were positioned in the interior-nanogap region. Both types of nanogap-less NPs (CAS NPs and gap-less AuNPs) showed similar increases in the SERS intensity. Thus, it can be concluded that it is the interior nanogap that ensures a highly enhanced SERS signal and that the composition of the metallic phase has only a minor effect on the SERS phenomenon. To examine the role of interior nanogap on SERS enhancement further, we placed the Raman reporter molecules (4-MPy) on the outmost shell surface instead of the Au core surface (Figure 3B). Irrespective of whether an interior nanogap was present or not, all similar-sized NPs (CAS NPs, gap-less AuNPs, and DIPs) showed a similar SERS signal, which was very weak. This result clearly suggests that the markedly stronger SERS enhancement was primarily attributable to the greater enhancement of the EM field in the interior nanogap.

Owing to the highly enhanced EM field and the uniformly confined hot spots in the interior nanogap, the SERS signal of the solution of the DIPs could be detected even at a very low concentration (500 fM), with there being a highly linear relationship ($R^2 = 0.99$) between the NP concentration and the SERS signal intensity (Figure 3C). In addition, we obtained stable SERS signals during time-course measurements performed under continuous laser exposure (Figure 3D). These experimental results indicate that the DIPs generated highly robust and sustainable SERS signals because of the Raman molecules stably encapsulated within highly SERS-active narrow interior nanogap. This property of the DIPs, wherein they produce uniform and quantitatively reproducible SERS signals with high sensitivity, can be exploited for biosensing/imaging. In addition, the DIPs were highly responsive under excitation at a wavelength of 785 nm (Figure S7); this suggests that they should be suitable for use for in/ex

![Figure 3. Nanostructure, Raman-dye position, particle concentration, and time-dependent SERS properties of as-synthesized NPs in solution.](image-url)
vivo and in vitro SERS imaging as well as photothermal therapeutic probes. On the other hand, the DIPs did not generate a detectable signal under excitation at a wavelength of 514 nm (Figure S7). Because the near-field enhancement of the SERS effect is determined by the plasmonic excitation mode near the resonance peak, we believe that the 514 nm laser, whose wavelength is much different from that corresponding to the near-field resonance peak induced by the plasmonic excitation mode of the interior nanogap, could not generate a sufficiently strong EM field in the interior-nanogap region.

**Single-Particle-Level SERS Analysis on DIPs.** To investigate the optical properties of the DIPs in more detail, we performed an analysis of the SERS spectrum of the DIPs at the single-particle level using an atomic force microscopy (AFM)-correlated nano-Raman instrument (Figure 4A). To acquire the SERS spectra of the individual DIPs, we first accurately matched the end of the AFM tip with the focal spot of the objective lens (see details in Supporting Information). Then, we simultaneously obtained the AFM topographical image and the Rayleigh scattering image (Figure 4B) from the same particle. The representative tapping-mode AFM image (10 × 10 μm²) indicated that the individual DIPs were well dispersed and did not overlap in the laser exposure focal spot (~250 nm). Furthermore, high-magnification AFM images confirmed the spherical shape of the DIPs, with their height profile matching well with their overall size as determined from the TEM images (Figure 4C). We measured the SERS spectra of 110 individual DIPs and calculated the enhancement factor (EF) values for the fingerprint peak at 1003 cm⁻¹ (see details in Supporting Information).

**Figure 4.** AFM-correlated Raman spectroscopy-based single-particle mapping analysis, SERS enhancement factor (EF) distribution, and polarization-resolved SERS plot for DIPs. (A) Instrumental setup used for AFM-correlated single-particle Raman spectroscopy. (B) Topographical matching of AFM image (left) and Rayleigh scattering image (right) for DIPs. (C) Magnified AFM image of a single DIP (left) and height profile across the NP (right, the red line in the AFM image). (D) A distribution of the SERS EF values at 1003 cm⁻¹ as measured from individual DIPs (110 particles were analyzed). The EF values show a narrow distribution of large SERS EFs, range from 1.1 × 10⁸ to 5.3 × 10⁹ for 90.0% and 97.3% of particle populations, respectively. All the analyzed particles displayed the EFs of 1.1 × 10⁸ or larger. (E) Single-particle polarization-resolved plot of the SERS intensity at 1097 cm⁻¹ with respect to rotation angle (θ). (F and G) The calculated polarization-resolved plots of the EM field in the interior-nanogap region with respect to a rotation angle for (F) DIP and (G) Au-NNP, respectively. The theoretical calculations were performed using the finite element method with COMSOL, and the maximal values of the EM field enhancement at each rotation angle were plotted. The black arrows in panel G indicate Au nanobridges.
structures with a nanogap can be exploited for use as reproducible and reliable SERS probes. It should be noted that the EF values of DIPs were conservatively underestimated by assuming the maximum number of 4-MPy molecules were uniformly packed on AuNP core surface (see details in Supporting Information). In addition, nonresonant Raman dyes (4-MPy) with a small cross-section were used in our study. The EF values of the DIPs are 2−3 orders of magnitude higher than those with other nanostructures reported previously for the same Raman dye molecule (4-MPy). It is now widely accepted that an EF of 10^6 to 10^8 could be sufficient even for single-molecule detection. In this respect, the as-synthesized DIPs, which generated very strong SERS signals with high sensitivity, are highly promising as SERS probes for use in various analytical applications. Significantly, we investigated the incident-laser-polarization-dependent SERS characteristics of the DIPs at the single-particle level (Figure 4E). In the case of anisotropic or assembled nanostructures, the optical properties are largely dependent on the geometric configuration and laser polarization direction. In contrast, the DIPs exhibited stable and uniform SERS signals regardless of the polarization direction of the incident laser, owing to the symmetrically distributed hot spots within the interior nanogap. This polarization-independent property also makes the DIPs suitable for use as highly reliable, quantitative analytical probes.

Figure 5. SERS-based ultrasensitive DNA detection assay using DNA-functionalized DIPs. (A) Schematic illustration of SERS-based ultrasensitive DNA detection assay with DNA-modified DIP nanoparticles that sandwich-capture target DNA with DNA-modified magnetic microparticles. (B and C) SEM images of target-DNA-specific hybridization complexes (target-captured DIP probe-MMP complexes) formed using different linker DNA [(B) complementary sequence DNA (HAV); (C) noncomplementary sequence DNA (HBV)]. Inset images: color of assay solution under external magnetic field after DNA sandwich hybridization. Scale bars are 500 nm. (D) SERS-based DNA detection assay results with DNA-modified DIPs (SERS intensities at 1003 cm⁻¹ were measured for different DNA concentrations of HAV (10 aM to 1 pM) and HBV (1 pM). Inset: Change in SERS spectra obtained with different concentrations of target DNA. All the spectra were obtained using 785 nm excitation laser at laser power of 2 mW and acquisition time of 5 s. SERS intensities were obtained and averaged with consecutive accumulation of five measurements for each concentration.

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DNA Assays with DIPs. To test the potential of DIPs as SERS sensing bioprobes, we performed SERS-based ultra-sensitive DNA detection assays using the NPs. In order to efficiently capture and separate target DNA, we used DNA-modified magnetic microparticles (DNA-MMPs) and DIP Raman probes (DIP probes) to detect the target DNA strands (hepatitis A virus DNA; HAV DNA), based on a typical sandwich-hybridization assay (Figure 5A, see details in Supporting Information). To confirm the formation of the sandwich hybridization complexes (target-captured DIP-MMP complexes) and its ability to undergo target-DNA-specific hybridization, we performed hybridization assays with complementary (HAV) and noncomplementary (HBV) DNA sequences (Figure 5A, see details in Supporting Information). In the presence of the complementary DNA, the solution turned colorless when an external magnetic field was applied to it, indicating the formation of the sandwich hybridization complexes (Figure 5B inset). In the presence of the noncomplementary DNA, however, the solution remained with the color even after the MMPs were collected with a magnet, meaning that the DIP probes were not captured by the DNA-MMPS (Figure 5C inset). Moreover, the formation of the sandwich hybridization complexes could be seen clearly in the scanning electron microscopy (SEM) images (Figure 5B, C). These results indicate that the proposed DNA detection method with DIPs is suitable for use in DNA detection assays. Figure 5D shows the changes in the SERS intensity at 1003 cm$^{-1}$ for varying concentrations of target DNA (HAV) solutions (10 aM to 1 pM, black squares). The SERS intensity quantitatively decreases with a decrease in the concentration of the target DNA over a wide range. On the other hand, with the intensity of the SERS signal was very low during control experiments in which the target DNA was replaced with noncomplementary DNA (HBV) strands (1 pM, red circular dot). The limit of detection (LOD) was approximatively 10$^{-10}$ aM, which is ~10 to 100-fold higher sensitivity than the DNA detection results with other nanostructures and methods. In terms of synthetic yield, structural precision, SERS EF distribution, and SERS signal stability and reproducibility, DIPs generated superior results over other SERS structures, resulting in highly sensitive and quantitative DNA detection. These results further confirm that the plasmonic DIPs fabricated with the SID process can be

Figure 6. SERS-based integrin ανβ3-specific cell imaging using peptide-functionalized DIPs. (A) Schematic illustration of SERS-based target-specific cell imaging with cRGD-modified DIP nanoparticles. (B and C) Bright-field microscopy images (left) and SERS maps (right) of U87MG cells (B, high integrin ανβ3 expression) and MCF-7 cells (C, integrin ανβ3 negative) incubated with cRGD-functionalized DIPs. Scale bar is 10 μm. SERS intensity at each mapping pixel (2 μm × 2 μm) was integrated for SERS spectra ranging from 983 cm$^{-1}$ to 1023 cm$^{-1}$ indicated by the shaded region in Figure 6D and color-scaled for cell imaging. All the spectra were obtained using 785 nm excitation laser at 4 mW power and with acquisition time of 1 s. (D) The SERS spectra obtained from the numerically marked positions in Figure 6B and 6C. (E) Time-dependent Raman profiles of cRGD-functionalized DIPs in a cell measured within the red box in panel B. (F and G) SERS maps of U87MG cells incubated with cRGD-functionalized (F) DIPs and (G, average particle diameter of 80 nm) AuNPs, respectively. Scale bar is 10 μm. The SERS intensity at each mapping pixel (2 μm × 2 μm) was integrated for the SERS spectra ranging from 983 cm$^{-1}$ to 1023 cm$^{-1}$ and color-scaled for cell imaging. All the spectra were obtained using 633 nm excitation laser at 400 μW and with acquisition time of 1 s.
used as ultrasensitive DNA sensing probes in target-selective, ultrasensitive, and quantitative analytical SERS probes.

**Targeted Cell Imaging with DIPs.** Lastly, we performed SERS-based target-specific cell imaging using surface-functionalized DIPs (Figure 6A). To evaluate the integrin-targeting specificity of the DIPs, cyclo(Arg-Gly-Asp-D-Phe-Lys) (c-RGDyK), hereafter denoted as cRGD) peptide, which specifically binds to integrin $\alpha_v\beta_3$ (overexpressed in metastatic and endothelial tumor cells), was attached onto the surfaces of the DIPs. Next, we introduced cRGD-functionalized DIPs into different cell lines, namely, U87MG (high integrin $\alpha_v\beta_3$ expression) and MCF-7 (integrin $\alpha_v\beta_3$ negative) (see details in Supporting Information). Owing to the high expression of integrin $\alpha_v\beta_3$ in the U87MG cells and their preferential binding affinity with respect to cRGD peptide, strong SERS signals were clearly observed in the integrin-$\alpha_v\beta_3$-positive U87MG cells, confirming the target-specific binding and SERS imaging capabilities of the cRGD-functionalized DIPs (Figure 6B,D). On the other hand, a SERS signal was barely detected in the case of the integrin-$\alpha_v\beta_3$-negative MCF-7 cells, confirming that the cRGD-functionalized DIPs showed good target selectivity for SERS-based cell imaging (Figure 6C,D). Importantly, the cRGD-functionalized DIPs generated uniform SERS signals steadily when they were continuously exposed by a laser for a long period (Figure 6E). DIPs produced more reliable, stronger SERS signals for cell imaging than AuNPs with a similar size (Figure 6F,G and Figure S8). It is worth noting that SERS-based cell imaging could be performed with a lower-power laser (400 $\mu$W) and short exposure time (1 s for each pixel) with DIPs, which are critical in minimizing cell and tissue damage and stably imaging cells for a long-term.

**CONCLUSIONS**

We developed a selective, interdiffusive dealloying-based facile synthetic strategy for forming highly SERS-active Au–Ag NPs (DIPs) containing a uniformly confined interior nanogap in a high yield (~95%). This strategy is a very simple, cheap, and efficient one for fabricating interlayer-free interior nanogap particles without using modulating interlayer materials, such as DNA, polymer, or silica shell. It was confirmed that the interior nanogap, as small as $\sim 2$ nm, was formed by selective Ag-etching and interdiffusion of Ag atoms from the Au–Ag alloy shell (SID process). It should be noted that metal residues were randomly distributed inside the gap. The NPs with the interior nanogap showed a highly enhanced SERS signal as compared to those of the shell-less or nanogap-less NPs with a similar size, owing to the strong EM field generated in the interior nanogap by strong plasmonic coupling between the core and the shell. Furthermore, the DIPs emitted robust, stable, and quantitatively reproducible SERS signals and exhibited a highly enhanced and narrowly distributed EFs, ranging from $1.1 \times 10^8$ to $5.3 \times 10^9$ for 97.3% of the particle population. Significantly, all the analyzed particles displayed the EFs of $\geq 1.1 \times 10^8$. We developed an ultrasensitive DNA detection assay with DIPs for 10 aM to 1 pM target concentrations that were clearly discernible with the SERS signals from DNA-modified DIPs. It was also shown the cRGD-functionalized DIPs can efficiently target integrin-overexpressed cells and generate uniform SERS signals steadily when they were continuously exposed by a laser for a long period. DIPs facilitated SERS cell imaging with a low-power laser (400 $\mu$W) and short exposure time (1 s for each pixel), and these allow for minimizing cell and tissue damages and stably imaging cells for a long-term (>30 min). Our synthetic strategy with the SID process offers new revenues in forming plasmonic nanostructures with a nanogap or strong plasmonic coupling without a need for an interlayer between two nanostructures (e.g., core and shell). DIPs show potential as strong, controllable, and quantitative SERS probes with a narrow distribution of high EFs without the dependence of a laser polarization direction, and can address the long-standing issues in SERS signal reproducibility, reliability, and quantification capability. Finally, DNA detection and cell imaging results with biofunctionalized DIP probes validate and provide the opportunities in utilizing SERS probes in sensitive, selective biosensing, bioimaging, and theranostic applications with a high reliability.1,6,21,43-45

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acscentsci.7b00584.

Experimental details, supplementary figures and table on synthesis and characterization of nanoparticles, Raman measurement, theoretical calculations, DNA assay, and targeted cell imaging (PDF)

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**Notes**

The authors declare no competing financial interest.

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