Repeated Solid-state Dewetting of Thin Gold Films for Nanogap-rich Plasmonic Nanoislands

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This work reports a facile wafer-level fabrication for nanogap-rich gold nanoislands for highly sensitive surface enhanced Raman scattering (SERS) by repeating solid-state thermal dewetting of thin gold film. The method provides enlarged gold nanoislands with small gap spacing, which increase the number of electromagnetic hotspots and thus enhance the extinction intensity as well as the tunability for plasmon resonance wavelength. The plasmonic nanoislands from repeated dewetting substantially increase SERS enhancement factor over one order-of-magnitude higher than those from a single-step dewetting process and they allow ultrasensitive SERS detection of a neurotransmitter with extremely low Raman activity. This simple method provides many opportunities for engineering plasmonics for ultrasensitive detection and highly efficient photon collection.

Spatial localization of intense electromagnetic (EM) fields near the nanostructures termed in 'hotspots' can be created by strong coupling of plasmonic near-fields between neighboring metallic nanostructures. The hot spots bring about the extraordinary increase of Raman scattering from neighboring molecules, i.e., surface enhanced Raman scattering (SERS). Recently, multiple hotspots between diverse plasmonic nanostructures advance the SERS detection of small biomolecules at low concentrations and even at a single-molecule level. However, this ultrasensitive detection still faces a problem with the reproducibility of plasmonic nanostructures for stable Raman signal enhancement and thus the construction of nanogap-rich SERS-active nanostructures still remains challenging. Based on the site enhancement in SERS, highly active SERS sites account for only 63 sites in 1,000,000 of the total sites but contribute to 24% of the overall SERS intensity.

High yield nanofabrication of nanogap-rich SERS substrates such as randomly dispersed nanoparticles, nanometer scaled rough film or well-defined nanostructures via e-beam lithography (EBL) or nanosphere lithography (NSL) and ion etching, has attracted substantial interest over the last decade. More recently, the combinations of nanolithographic patterning with templated growth or high aspect ratio negative resist process enable the nanofabrication of sub-nanogap metal nanostructures whereas these methods still have technical challenges such as low cost, large area fabrication, and tuning capability of plasmon resonance wavelength. Lithography-free nanofabrications by using self-assembled metal nanoparticles in solution phase can also overcome such limitations with large-area homogeneity in a highly reproducible manner. However, additional chemical modification and other driving forces such as electrostatic and hydrodynamic interaction are still required for assembling metallic nanoparticles on a substrate. Alternatively, solid-state dewetting of metal thin films has distinctive advantages for producing metal nanoisland arrays over a large area with enhancement factor over $10^5$. The average size and separation of metal nanoislands can be moderately controlled with the initial thickness of a thin metal film. However, the concurrent fabrication of small gap spacing and large nanoisland size still remains challenging: as the film thickness increases, both the average nanoisland size and the

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interstitial gap spacing simultaneously increase after a single-step solid-state dewetting process (see Fig. S1 in supplementary information). This unique nature intrinsically confines the use of solid-state metal dewetting in fabricating highly scattered SERS substrates, i.e., large nanoislands with small gap spacing. Moreover, this trade-off between the size and the gap hinders the precise tuning of plasmon resonance wavelength because the plasmon resonance typically becomes red-shifted as the nanoisland size increases but blue-shifted as the gap spacing increases\(^{22,23}\). Very recently, multiple dewetting allows the size control of nanoislands\(^{24}\), however, the quantitative analysis is still in need for the size-gap control and plasmon resonance tuning as a practical guideline for developing engineered plasmonic nanostructures.

Here, we report nanogap-rich gold nanoislands with enlarged scales for highly sensitive SERS substrates by using simply repeating solid-state dewetting of thin Au film. The schematic illustration compares two different Au nanoislands created by using a single and repeated solid-state dewetting (Fig. 1a,b). Unlike a single-step dewetting, the repeated dewetting can provide enlarged Au nanoislands with small gap spacing, which increase the number of plasmonic hotspots and thus enhance the extinction intensity as well as the tunability for plasmon resonance wavelength. The plasmon resonance was numerically calculated by using a finite-difference time-domain (FDTD) method, where the Au nanoisland was modeled as nanoisland arrays in a square lattice with a period of 100 nm. As the nanoisland diameter increases, the gap spacing between neighboring nanoislands decreases simultaneously. The calculated results clearly indicate that a substantial increase of the hotspots owing to the enlarged size with small gap spacing apparently contributes to an increase of the extinction intensity as well as a red-shift of the plasmon resonance wavelength. (Fig. 1c).

**Results and Discussion**

Nanogap-rich Au nanoislands were simply achieved by alternating thin Au film evaporation and thermal annealing. In the first step, a thin Au film was thermally evaporated on a 4 inch quartz substrate and then thermally annealed to form Au nanoislands on a hot plate for one hour at 500 °C. In the second step, the second thin Au film was thermally evaporated on the pre-determined Au nanoislands and subsequently annealed at the same condition (Fig. 2a). This second step not only reduces the gap spacing between neighboring Au nanoislands but also increases the nanoisland size. A thin Au film on the Volmer-Weber mode\(^{25}\) of thin film growth initially forms isolated nanoislands, nucleates, and coalesces each other during the first step. However, the pre-determined nanoislands during the second step accelerates the rapid coalescence of nanoislands rather than the nucleation. Consequently, the second dewetting process enlarges the size of initially formed Au nanoislands as well as reduces the interstitial gap spacing between neighboring Au nanoislands. In this experiment, the repeated solid-state dewetting was consecutively performed from thermal evaporation and dewetting of additional 5 nm thick Au film after thermal dewetting of Au film with 5 nm in initial thickness (denoted by \(t_{1Au} = 5 \text{ nm}, t_{2Au} = 5 \text{ nm}\)). For
comparison, Au nanoislands after a single-step was also obtained by using Au film with 5 nm (denoted by $t_{1\text{Au}} = 5 \text{ nm}$) and 10 nm (denoted by $t_{1\text{Au}} = 10 \text{ nm}$) in initial thickness, respectively. The SEM and AFM images show Au nanoislands under different dewetting conditions, i.e., $t_{1\text{Au}} = 5 \text{ nm}$, $t_{1\text{Au}} = 10 \text{ nm}$, $t_{2\text{Au}} = 5 \text{ nm}$ and $t_{1\text{Au}} = 10 \text{ nm}$, shown in Fig. 2b. Both the microscopic images apparently demonstrate enlarged Au nanoislands with small gap spacing after the repeating solid-state dewetting of thin Au film, compared to those after the single-step. Besides, the side view TEM images of Au nanoislands clearly support the enlarged size with small gap spacing (Fig. 2c). The effective diameter and packing density of Au nanoislands were calculated from the binary segmentation of Au nanoislands shown in the SEM images, where the error bar indicates the size distribution of Au nanoislands (Fig. 2d). This repeated solid-state dewetting process clearly exhibits both the high packing density from the enlarged effective diameter with small gap spacing and the relatively small size distribution, compared to a single-step solid-state dewetting, i.e., Au nanoislands from Au film with 5 nm ($t_{1\text{Au}} = 5 \text{ nm}$) and 10 nm ($t_{1\text{Au}} = 10 \text{ nm}$) in initial thickness. The optical images also demonstrate a wafer-level fabrication of enlarged Au nanoislands with small gap spacing (Fig. 2e).

The enlarged Au nanoislands with small gap spacing apparently increase the number of plasmonic hotspots and thus enhance the extinction intensity as well as the tunability for plasmon resonance wavelength. The size and interstitial gap spacing of pre-determined Au nanoislands can be moderately controlled by the initial Au film thickness. The effective diameters of Au nanoislands from 5, 7, and 9 nm in initial thickness during the first dewetting process are about 25 nm, 31 nm, and 41 nm, respectively. In the first dewetting process, the gap spacing between neighboring Au nanoislands inherently increases with the nanoisland size, which also increases with the initial film thickness. In particular, an increase of the film thickness reduces the nucleated void density and widens the void opening during dewetting to reach a stable configurations, which consequently increases the gap spacing between Au nanoislands. However, the repeated solid-state dewetting significantly suppresses an increase of the gap spacing whereas the nanoisland size becomes enlarged. This geometrical variation of Au nanoislands favorably contributes to a substantial red-shift of the plasmon resonance wavelength (PRW). For different Au film thicknesses of 5 nm, 7 nm, and 9 nm, the extinction spectra from a single-step (black line) and repeated dewetting (red line) clearly demonstrate that enlarged nanoislands with small gap spacing after repeated dewetting enhances both the extinction intensity and the red-shift of PRW (Fig. 3a). The experimental results also include the binary segments of Au nanoislands from the SEM images (left).
and the electric field intensity distributions at their corresponding plasmon resonance wavelengths were numerically calculated by using a finite-difference time-domain (FDTD) method (right) (see Fig. S2. in supplementary information). A decrease in gap spacing results in stronger electromagnetic hotspots. The effective diameter and gap of Au nanoislands after a single-step and repeated dewetting were calculated from the binary segmented SEM images for different film thicknesses (Fig. 3b,c, Fig. S3 in Supplementary Information). The results clearly exhibit that the nanoisland size increases by more than twice but the gap spacing relatively remains constant after the repeated dewetting with double film thickness for different film thicknesses. In addition, based on the comparison between a single-step (dashed line) and repeated (solid line) solid-state dewetting, the effective diameter increases but the gap spacing remain constant as the total Au film thickness increases and therefore it concludes that the repeated dewetting enables the fabrication of small gap spacing Au nanoislands (Fig. 3d). Depending on the effective diameter, the calculated PRW (Fig. S4 in supplementary information) and normalized extinction intensity well agree with the experimental results, which indicate that both the PRW shift and the extinction values increases with the effective diameter (Fig. 3e,f). If a given Au thickness for the first dewetting process is thicker than additional film for the second dewetting process, the thinner additional film for the second dewetting process may give rise to multiscale nanoislands (Fig. S5 in supplementary information). This scaling behavior of the plasmon resonance coupling between Au nanoislands is because the plasmon coupling strength varies universally as a function of the size and gap spacing. Thus the repeated dewetting enables precise tuning of the PRW over a wide wavelength range by changing the size and gap spacing.

Highly dense EM hotspots and PRW matching of Au nanoislands are very crucial in achieving enhanced Raman signals from the plasmonic nanostructures23. In this experiments, SERS signals of benzenethiol (BT) molecules under an excitation laser of 633 nm were measured from two different Au

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**Figure 3.** Tuning capabilities of Au nanoislands by using the repeated dewetting process. (a) Extinction spectra of a single-step and repeated solid-state dewetting of thin Au film with (i) 5 nm, (ii) 7 nm, and (iii) 9 nm in thickness. Binary segments of Au nanoislands from the SEM images (left) and numerically calculated electric field intensity distributions at the plasmon resonance wavelength from the SEM images by using a three-dimensional finite difference time domain (3D-FDTD) method (right). (b) the effective diameters and (c) the effective gaps after a single-step and repeated solid-state dewetting processes. (d) the effective diameter and the gap depending on total Au film thickness. The black and blue lines indicate a single-step and repeated dewetting, respectively. The 3D-FDTD calculated and experimental (e) plasmon resonance wavelengths and (f) corresponding normalized extinctions of Au nanoislands after a single-step and double dewetting, i.e., repeated dewetting with the same thickness, depending on the effective diameter. The black and red lines indicate a single-step and repeated dewetting, respectively.
nanoislands with a same total Au film thickness, i.e., Au nanoislands after a single-step dewetting of 10 nm in initial thickness (\(t_{1Au} = 10\) nm) and after repeated dewetting with 5 nm in both initial and additional thickness (\(t_{1Au} = 5\) nm, \(t_{2Au} = 5\) nm), respectively. The result shows the SEM images, the calculated electric field intensity distributions at an excitation wavelength of 633 nm (upper left), and representative SERS spectra of BT molecules (upper right) (Fig. 4a,b). In this experiment, the SERS signals at 1069 cm\(^{-1}\) of BT molecules were quantitatively measured under 633 nm excitation and the corresponding SERS enhancement factor (EF), i.e., the average signal enhancement from absorbed molecules participated in SERS to that in Raman scattering, was spatially mapped with 1 cm intervals over two different 4 inch substrates. An expanded spatial map of SERS EFs at 1069 cm\(^{-1}\) were acquired at 1 mm intervals over an area of 1 cm\(^2\) and has spatially averaged SERS EFs about \(10^5\) for 10 nm and \(10^7\) for \(t_{1Au} = 5\) and \(t_{2Au} = 5\) nm, demonstrating the excellent uniformity over a 4-inch substrate. The SEM images and numerically calculated electric field intensity distributions at excitation wavelength of 633 nm by using a three-dimensional finite difference time domain (3D-FDTD) method (upper left) and representative SERS spectra of BT molecules (upper right). (c-d) SERS limit-of-detection (LOD) of octopamine molecules with different concentrations. The LOD exhibits ca. 100\(\mu\)M for (c) a single-step solid-state dewetting with \(t_{1Au} = 10\) nm but increases down to 10\(\mu\)M for (d) repeated dewetting with \(t_{1Au} = 5\), \(t_{2Au} = 5\) nm.

Figure 4. SERS enhancement factors (EF) of Au nanoislands and their SERS limit-of-detection. a,b) Spatial distribution of SERS enhancement factors (EFs) at \(1069\) cm\(^{-1}\) of benzenethiol (BT) molecules at an excitation laser of 633 nm over an area of 4-inch glass substrate for (a) a single-step solid-state dewetting with \(t_{1Au} = 10\) nm and (b) a repeated dewetting with \(t_{1Au} = 5\), \(t_{2Au} = 5\) nm. A expanded spatial map of SERS EFs at \(1069\) cm\(^{-1}\) were acquired at 1 mm intervals over an area of 1 cm\(^2\) and has spatially averaged SERS EFs about \(10^5\) for 10 nm and \(10^7\) for \(t_{1Au} = 5\) and \(t_{2Au} = 5\) nm, demonstrating the excellent uniformity over a 4-inch substrate. The SEM images and numerically calculated electric field intensity distributions at excitation wavelength of 633 nm by using a three-dimensional finite difference time domain (3D-FDTD) method (upper left) and representative SERS spectra of BT molecules (upper right). (c-d) SERS limit-of-detection (LOD) of octopamine molecules with different concentrations. The LOD exhibits ca. 100\(\mu\)M for (c) a single-step solid-state dewetting with \(t_{1Au} = 10\) nm but increases down to 10\(\mu\)M for (d) repeated dewetting with \(t_{1Au} = 5\), \(t_{2Au} = 5\) nm.
The dewetting process, however, the LOD for repeated dewetting is increased by one order of magnitude, i.e., ca. 10 μM.

To be concluded, this work reports a simple and wafer-level fabrication for nanogap-rich metal nanoislands with enlarged sizes for highly sensitive SERS substrates by repeated solid-state dewetting of thin Au film. The precise control of the initial and additional Au film thicknesses enables manipulation of the size, interstitial gap spacing, packing density and plasmonic properties of Au nanoislands. The experimental results clearly demonstrate that Au nanoislands from repeated dewetting increases the SERS EF by about 7.5 times, compared to those from the single-step dewetting process. This repeated dewetting process can provide many opportunities for not only engineering plasmonics for ultrasensitive detection and highly efficient photon collection but also for nanoscale metal masks for reactive ion etching or seeds for the catalytic growth of nanowires.

Methods

Extinction measurement. The extinction spectra were measured with a microscopic spectrometer, which consists of an inverted microscope (Carl Zeiss Axiovert 200M) equipped with an excitation laser, and a spectrometer (MicroSpec 2300i) with a charge-coupled device (CCD) camera (Princeton Instruments, Model PIXIS: 400BR). The extinction spectra of silver nanoislands were collected with 50× lens (NA = 0.5).

SERS measurement. A helium–neon laser (632.8 nm) and a spectrometer (MicroSpec 2300i spectrometer equipped with a charge-coupled device camera (Model PIXIS: 400BR, Princeton Instruments)) were coupled with an inverted microscope (Axiovert 200M, Zeiss). The excitation and collection of light were done by a 50× objective lens (NA 0.5). The excitation laser power was 5.15 mW and the acquisition time was 1 second.

SERS enhancement factor (EF). SERS EF was calculated by the conventional method, where the average enhancement factor of $E_{F\text{ave}}$ is defined as $E_{F\text{ave}} = \frac{I_{\text{SERS}}/N_{\text{SERS}}}{I_{\text{Raman}}/N_{\text{Raman}}}$, where $I_{\text{SERS}}$ is the SERS intensity, $I_{\text{Raman}}$ is the Raman intensity, $N_{\text{SERS}}$ is the number of probed molecules in SERS, and $N_{\text{Raman}}$ is the number of probed molecules in Raman measurements. The intensity of the 1069 cm$^{-1}$ peak of BT molecules from the SERS and Raman measurements have been used for SERS EF calculation. The effective SERS active surface area of repeated dewetting from Au film of 5 nm in both initial and additional thicknesses is increased by ca. 2.2 times, compared to that from a single-step dewetting of 10 nm thick Au film. The surface coverage of benzenthiol self-assembled on gold is about 6.8×10$^{14}$/cm$^2$. The extinction of BT molecules is 1.3×10$^{13}$ and the Raman intensity ($I_{\text{Raman}}$) is 3.7×10$^{6}$ for a single-step of solid-state dewetting of 10 nm thick Au film and 7.4×10$^{7}$ for repeated dewetting from Au film of 5 nm in both initial and additional thicknesses. The number of BT molecules ($N_{\text{Raman}}$) was calculated from the average number of molecules in the detection volume. The detection beam spot radius is about 2.5 μm and experimentally measured collection depth is about 110 μm. The density of BT molecules is 1.073 g/ml, and the molecular weight is 110.18 g/mol. Therefore, the number of BT molecules ($N_{\text{Raman}}$) contributing to the normal Raman signal measured from the standard is 1.3×10$^{13}$ and the Raman intensity ($I_{\text{Raman}}$) over the number of probed molecules in Raman measurements ($N_{\text{Raman}}$) is 3.7×10$^{13}$. The calculation results described above, $E_{F\text{ave}}$ is about 2.7×10$^{4}$ for a single-step dewetting of 10 nm thick Au film and 2.0×10$^{7}$ for repeated dewetting from Au film of 5 nm in both initial and additional thicknesses.

Numerical analysis of E-field distribution. The electric fields of Au nanoislands from different initial and additional thicknesses were numerically calculated by using a three-dimensional finite-difference time-domain method (Numerical FDTD Solutions 8.0.6). The geometric configuration of Au nanoislands were extracted from the binary segmented SEM images by using Image J software and the individual Au nanoislands were considered as a cylindrical shape (see Fig. S2 in supplementary information).

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Additional Information
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