Gold nanoparticle transfer through photothermal effects in a metamaterial absorber by nanosecond laser

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A non-complicated, controllable method of metallic nanoparticle fabrication at low operating light power is proposed. The method is based on laser-induced forward transfer, using a metamaterial absorber as the donor to significantly enhance the photothermal effect and reduce the operating light fluence to 35 mJ/cm², which is much lower than that in previous works. A large number of metallic nanoparticles can be transferred by one shot of focused nanosecond laser pulses. Transferred nanoparticles exhibit good size uniformity and the sizes are controllable. The optical properties of transferred particles are characterized by dark-field spectroscopy and the experimental results agree with the simulation results.

Fabrication of metallic nanostructures has drawn much attention in recent years because of its importance in applications in various fields such as optics, chemistry, microelectronics, biomedicine, etc.1–9. The laser-induced forward transfer (LIFT), as one of the methods, has been extensively studied due to its ability to deposit nanostructures of diverse materials onto different receiver substrates10–24. However, in previous works, femtosecond laser is typically utilised10–20, while only a few of them are realised by nanosecond laser pulses21–24. Due to the high melting point of noble metals that are commonly used (bulk Au 1337K, bulk Ag 1235K, etc.), high operating light power is also required. The reported threshold light fluences for metal transferring mostly varied from about 100 mJ/cm² to several J/cm²11–23, while for those with fluence under 100 mJ/cm², fs-pulse laser is utilised10. Furthermore, only one nanoparticle (NP)12–21 or a few NPs without regular morphologies11 are transferred by one light shot in most previous reports. Large-scale transferring methods are rarely studied10,23. In their works, fs-pulse lasers10 or ns-pulse lasers with large fluence (460 mJ/cm²)23 are needed, respectively.

In this article we report a non-complicated method of large-scale gold NPs fabrication through nanosecond laser pulses at low operating fluence based on LIFT. A metamaterial absorber sample is utilised as the donor substrate. The absorber performs nearly perfect absorption at specific wavelength25,26, thus significantly enhancing the photothermal effect around the exposed area. Therefore, the transferring can be realised by ns-pulse lasers with the threshold light fluence reduced to only 35 mJ/cm². A large number of NPs are transferred by only one light shot. These NPs show good size uniformity and the sizes can be pre-designed due to the mass conservation during the transferring process. Further studies of the transferred NPs are conducted through dark-field spectroscopy, scanning electron microscope (SEM) and transmission electron microscope (TEM). It is found that the properties are comparable to those of Au NPs prepared by chemical methods, as can be confirmed by our numerical simulations.

Experimental section

In our experiment, a metamaterial absorber is used as the donor substrate in order to utilise its high absorption to enhance the photothermal effect, thus significantly reducing the operating light fluence. The metamaterial absorber is a three-layer metal-insulator-metal structure, consisting of periodic Au nano-disk (ND) arrays (top) and a continuous Au film (bottom) separated by a Al2O3 layer, as shown in the inset of Fig. 1. Thicknesses of the three layers from top to bottom are 30 nm, 18 nm and 80 nm, respectively. The absorber displays a high-efficient absorption for a specific wavelength25,26, thus significantly enhancing the photothermal effect around the exposed area. Therefore, the transferring can be realised by ns-pulse lasers with the threshold light fluence reduced to only 35 mJ/cm². A large number of NPs are transferred by only one light shot. These NPs show good size uniformity and the sizes can be pre-designed due to the mass conservation during the transferring process. Further studies of the transferred NPs are conducted through dark-field spectroscopy, scanning electron microscope (SEM) and transmission electron microscope (TEM). It is found that the properties are comparable to those of Au NPs prepared by chemical methods, as can be confirmed by our numerical simulations.
As shown in Fig. 1, the absorber sample is placed upside down with the receiver substrate directly attached underneath. The light source is a super continuum laser (NKT SuperK Compact) with operating wavelengths from 500 nm to 2400 nm, a single-pulse energy of 2 μJ and a pulse width of 3 ns. The laser beam, which is normally irradiated on the front side of the absorber sample through the receiver, is collimated and then focused by a 100× objective lens (Mitutoyo, M Plan Apo NIR), with a beam width of about 3 μm.

Due to the high optical absorption by the metamaterial absorber, abundant heat accumulates around the exposed area, resulting in temperature rising above the melting point of Au NDs. Therefore, the Au NDs in this area are melted and then accumulate around the exposed area, resulting in temperature rising above the melting point of Au NDs. Consequently, the laser-induced nanoparticle transfer is realized. Owing to the absorber’s high-efficient absorption, the operating light fluence is greatly reduced to only 35 mJ/cm².

Results
The absorber sample after transferring is characterized under SEM, as shown in Fig. 2. A clear circle is created by one shot, and the diameter of the circle is roughly the same as that of the light beam. The gold NDs inside the circle are transferred. Outside the circle, the light power is not high enough to melt the NDs, so they remain the original disk morphology. Additionally, around the edge light power is not high enough to melt the NDs, so they remain the original disk morphology. The transferred nano-droplets cool down and solidify again into spherical dome shapes. Consequently, the laser-induced nanoparticle transfer is realized. Owing to the absorber’s high-efficient absorption, the operating light fluence is greatly reduced to only 35 mJ/cm².

The TEM image of the re-solidified Au NPs shows a lattice fringe spacing of about 0.24 nm (Fig. 3(d)), indicating the Au crystalline structure with (111) orientation. Additionally, energy spectrum of the transferred NPs is measured under SEM experiments, and the results verify that the particles are gold.

Discussion
Given the fact that the duration of one laser pulse is only nanosecond and the light power irradiated on the absorber sample is relatively low, gold sublimation does not occur and mass conservation of gold can be assumed. For a single transferred NP with a diameter of about 132 nm and diameter of 145 nm, the calculated volume is about 1.59 × 10⁻³ μm³, just slightly smaller than that (1.71 × 10⁻³ μm³) of the Au ND with a 270 nm diameter before transferring. It can also be found that the transferred NPs appear rather bigger than others. Take these particles labeled as 1 and 2 in Fig. 3(a) for example, their diameters are 212 nm, 232 nm and 251 nm, respectively. The corresponding volumes are about 4.87 × 10⁻³ μm³, 6.38 × 10⁻³ μm³ and 8.08 × 10⁻³ μm³, which are nearly 3, 4 and 5 times of a single particle, respectively. This is due to the fact that several melted gold NPs can merge together and then solidify into a larger sphere during the process of transferring. According to these calculations, mass conservation is satisfied during the transferring process and this ensures that the transferred NPs’ sizes can be controlled by designing the parameters of the original NDs. The two related varied parameters of NDs are the diameter and the period. The sizes of transferred NPs are mostly defined by the diameters of NDs, while periods only affect the distribution density of transferred NPs, which leads to the simplicity in the size controlling.

The TEM image of the re-solidified Au NPs shows a lattice fringe spacing of about 0.24 nm (Fig. 3(d)), indicating the Au crystalline structure with (111) orientation. Additionally, energy spectrum of the transferred NPs is measured under SEM experiments, and the results verify that the particles are gold.

The scattering spectra of single Au NPs are measured by a sensitive spectrometer (Ocean Optics QE65Pro). Here five samples with different transferred NP sizes transferred from five different samples are measured. Measuring spectra and corresponding SEM images are shown in Fig. 4. The diameters of the gold NPs are 86 nm, 115 nm, 135 nm, 148 nm and 172 nm, transferred from NDs with diameters of 120 nm, 180 nm, 230 nm, 265 nm and 330 nm, respectively. As the NP size increases, an obvious red-shift is observed from the measured scattering spectra (red dots), consistent with the simu-
The transferred NPs are randomly distributed on the receiver, and thereby two NPs located extremely close to each other form a dimer-like structure. The scattering spectra of these dimers excited by unpolarized incident light are measured. The dimers show some peculiar properties as compared with a single particle. For a dimer with a ~10 nm spacing between its two singles, its spectrum (red dots and lines) displays a distinct red-shift compared to that of the single NP (blue dots and lines), as demonstrated in Fig. 5(a). When the two singles forming the dimer are not in the same plane, as is shown in the inset of Fig. 5(b), two peaks can be observed in the scattering spectrum. All the experimental results agree well with the simulations.

Figure 3 | The transferred NPs are studied through dark-field spectroscopy, SEM and TEM. (a) SEM image of the receiver substrate; inset (i) is a 60° oblique viewing of a single transferred NP; inset (ii) is the schematic of a single transferred NP, exhibiting a spherical dome shape; here $h$ represents the height of the NP. (b) Histogram of the transferred NP size distribution. (c) Dark-field optical microscopy image; inset is the scattering spectrum of the NPs, and the area measured is shown by the green circle. (d) A transferred Au NP under TEM.

Figure 4 | Measured (red dots) and simulated (blue lines) scattering spectra and corresponding SEM images of transferred NPs with different sizes. The diameters are (a) 86 nm, (b) 115 nm, (c) 135 nm, (d) 148 nm and (e) 172 nm, respectively. A distinct red-shift can be observed as the NP size increases.
operating at low light fluence (35 mJ/cm²) is required. Owing to the process can be simply conducted in air, and only a ns-pulse laser proposed to mass-fabricate metal NPs through nanosecond laser simulation (red line) results. Furthermore, since mass conservation is satisfied, the transferred NPs' sizes are designable through controlling the original NDs' parameters. The optical properties of transferred single NPs and dimers are measured by dark-field spectroscopy, and the results are consistent with the simulations. Our method is therefore promising in nanostructures preparation for potential optoelectronics and biology applications, in particular, when diameters of NPs is required in advance, while with good uniformity.

Figure 5 | Scattering spectra of dimers are measured. (a) Scattering spectra of a dimer (red) and a single NP (blue) are measured (dots) and simulated (lines); and (b) for a dimer with its two singles not in the same plane, two peaks are observed according to the measured (red dots) and simulation (red line) results.

In conclusion, a novel, non-complicated method based on LIFT is proposed to mass-fabricate metal NPs through nanosecond laser pulses at low operating fluence. A metamaterial absorber is utilised as the donor to enhance the photothermal effect. The transferring process can be simply conducted in air, and only a ns-pulse laser operating at low light fluence (35 mJ/cm²) is required. Owing to the high density of NDs on absorber sample, massive NPs can be transferred within one irradiation, and the transferred NPs exhibit excellent size uniformity. Furthermore, since mass conservation is satisfied, the transferred NPs' sizes are designable through controlling the original NDs' parameters. The optical properties of transferred single NPs and dimers are measured by dark-field spectroscopy, and the results are consistent with the simulations. Our method is therefore promising in nanostructures preparation for potential optoelectronics and biology applications, in particular, when diameters of NPs is required in advance, while with good uniformity.

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Author contributions

MQ conceived the experiment, and supervised the entire project with Q.L. H.M.G. built the experimental setup with the help of X.C. and carried out the characterization with X.X.C. and D.Z. Y.Q.Y. conducted the numerical simulations and analyzed the data with H.M.G. Y.T.C. and M.Y. fabricated the samples. All the authors provided technical and scientific insight and contributed to the writing of the manuscript.

Additional information

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