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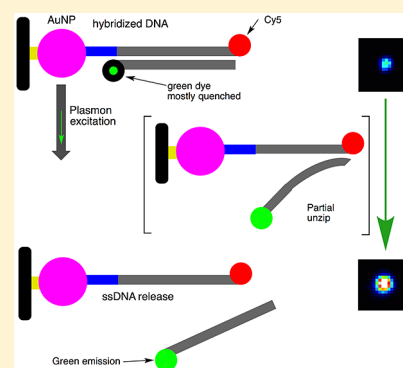
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Thermoplasmonic ssDNA Dynamic Release from Gold Nanoparticles Examined with Advanced Fluorescence Microscopy

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S Supporting Information

ABSTRACT: Plasmon excitation of spherical gold nanoparticles carrying a fluorescent labeled 30 bp dsDNA cargo, with one chain covalently attached through two S–Au bonds to the surface, results in release of the complementary strand as ssDNA that can be examined in situ using high-resolution fluorescence microscopy. The release is dependent on the total energy delivered, but not the rate of delivery, an important property for plasmonic applications in medicine, sensors, and plasmon-induced PCR.



Plasmon-mediated heating involves localized ultrafast energy-to-heat conversion upon light interaction with electrons of a given metal nanoparticle.¹ In the case of gold nanoparticles (AuNP), this heat can increase the temperature hundreds of degrees near the surface of the nanoparticle, for submicrosecond times following laser excitation.² However, thermoplasmonic effects in long time scales are determined by the principles of thermodynamics and the dynamics of heat dissipation¹ and have been recently utilized for compact autoclaves³ or in photothermal imaging for single particle detection;⁴ at very short time scales, molecules can experience high-temperature behavior that reflects their location and interaction with the plasmonic field.

Realistic technological applications of plasmon heating in devices based on nanostructures will require enabling chemical platforms to support the efficient energy delivery without affecting the surface of the nanoparticle. Here, we present a new platform; an *on–off* fluorescence switch is used to report single strand DNA (ssDNA) release from AuNP, where only one of the chains of double strand DNA (dsDNA) is covalently attached to the nanoparticle. Meanwhile, the second chain, attached only through DNA hybridization, is released upon plasmonic excitation. Beyond sensors⁵ and therapeutic applications,^{6,7} we foresee a potential for polymerase chain reaction (PCR) applications with a dramatic reduction of bulk heating requirements. Because thermoplasmonic strategies can deliver energy with pinpoint precision while the bulk solution remains at near ambient temperature, this will allow the developments of novel applications such as remote controlled

in vitro PCR. In the case of potential photoplasmonic PCR applications, a strategy that allows reversible binding to the AuNP surface would have to be developed; at the same time, the complexity of a design around available fluorescent dyes and laser wavelengths would not be required.

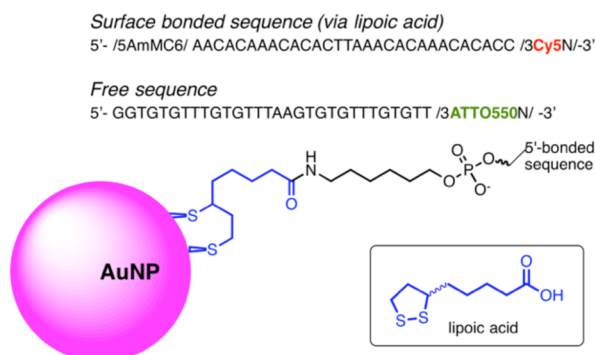
Branda et al.^{8–10} and Halas et al.^{11–13} have pioneered the field of plasmon-mediated release. Reports include the release of ssDNA with a complementary strand bound to a gold nanoparticle in solution.^{10,11,14} In Branda's approach, they linked their complementary 15bp strand through a 5' terminal –SH anchor¹⁵ to the nanoparticle surface. A threshold of ≈ 0.5 W/cm² was found as the limit for breaking the S–Au bond. Plasmon-induced release was also observed by Halas et al.^{11,13} One example has also been reported where local plasmonic heating hinders hybridization.¹⁶ In the present Letter, we examine the dynamics of ssDNA plasmon release by using an anchoring system composed of lipoic acid (LA), a molecule that has been employed as oligo-linker for gold nanoparticles,^{17,18} and a five-carbon spacer to a 30 bp duplex, see Scheme 1. We note that in this case ssDNA release does not involve covalent bond cleavage; further, the use of LA provides two S–Au links minimizing the risk of cleavage at the AuNP surface. Single strand release from the nanoparticle surface was in situ monitored for the first time using high-resolution

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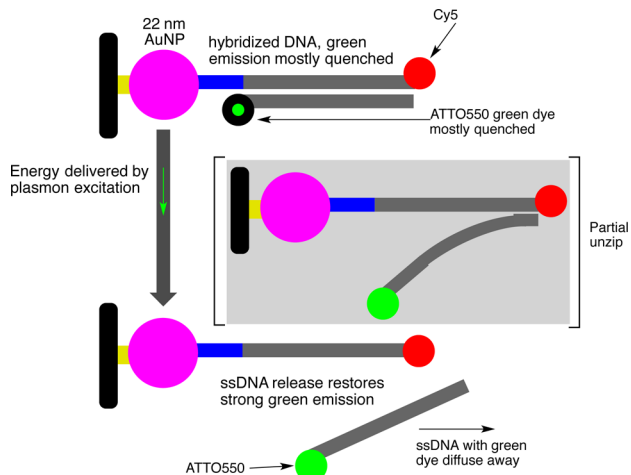
Scheme 1. Details of the Sequences Employed and Nature of the Lipoic Binding to AuNP



microscopy resolving emissive events and with spatial resolution at subcellular dimensions.

Scheme 2 shows a representation of the experimental design. 3-Aminopropyltriethoxysilane (APTES) was used to attach the

Scheme 2. Representation of dsDNA-AuNP Conjugate System^a



^adsDNA is attached through an amide bond between the bonded strand, Cy5 labeled, and the lipoic acid (in blue) functionalized AuNP. On the left, the black bar symbolizes the glass surface and APTES attachment (yellow). Upon plasmon excitation release of the ssDNA label with ATTO550 takes place with a concomitant increase in the fluorescence intensity (see Supporting Information Section S3). Covalent bond cleavage is not involved in the release process. The square brackets show a partially unzipped system, a necessary step in the release dynamics.

spherical 22 ± 3 nm AuNP onto the glass surface (see Supporting Information Section S2 and Figures S1 and S2). LA anchors through its two-thiol functionalities to the gold surface, Scheme 1; the “two-feet” surface binding prevents the release of dsDNA (with the Cy5 dye as a control, see Figure 1). Further, LA also serves as a spacer and anchor through its carboxylic moiety to the 5' amine strand in the 30 bp dsDNA shown in Scheme 1. The dsDNA was chemically attached to the gold surface, ranging between 200 and 500 dsDNA per nanoparticle (see Supporting Information), where the uncertainty reflects the allowance for the spacer that effectively provides some relief for DNA crowding on the surface. These characteristics are critical for our real-time fluorescence

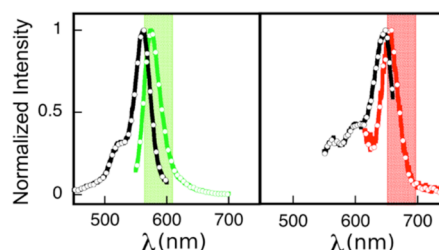


Figure 1. Excitation (black line) and emission (colored lines) spectra of the dsDNA labeled with ATTO550 (green, left panel) and Cy5 (red, right panel). The color-shaded areas represent the bandpass filters for the fluorescence emission.

detection. ATTO550, which is located at ~ 2.0 nm from the AuNP surface in the bonded oligo strand would be largely quenched and retain only 5.0–20% of its original brightness. Meanwhile, for Cy5, which is located ~ 8.0 nm from the metal surface would show 70–90% of its original brightness (see Supporting Information Section S3 and Figures S3 and S4). Our strategy uses Cy5 as a reporter to quantify loss of DNA binding due to potential, but highly unlikely, cleavage of the LA-AuNP linkage; see Scheme 2.

Total internal reflection fluorescence (TIRF) images, obtained with 543 or 633 nm CW laser excitation, for the samples containing the dsDNA attached to the AuNP were acquired. Image sequences taken up to 250 s using 543 nm excitation showed regions with intense emissions, whose intensities and size change as a function of the exposure time and energy (see Figure 2A for intensity changes for representative bright spots over the AuNP functionalized slide). Further, emission spectra and lifetimes for the fluorescent spots matched those of ATTO550 in solution as seen in Figure 2B. Experiments were performed at 633 nm to test Cy5 photostability, only produced bleaching of the AuNP-bound ssDNA Cy5 emission to a similar extent to that observed for the free construct in solution (see Supporting Information Figures S5 and S6).

The increase in signal intensity of ATTO550 can be interpreted as the specific release of the ATTO-labeled (i.e., the noncovalently bonded) strand from the quenching zone of the AuNP as depicted in Scheme 2. Physical desorption from the gold surface can be ruled out as an important factor responsible for the ATTO550 emission changes because experiments with noncomplementary ATTO550 labeled oligonucleotides or free ATTO550 dye showed no emission changes (data not shown). To further verify that the release of noncovalently bonded ssDNA takes place and not of the bonded dsDNA, control experiments were carried out. First, a 530 nm LED as an illumination source (0.32 W/cm^2) was used over the dsDNA@AuNP slides, revealing only the presence of ATTO550 fluorescence with nondetectable Cy5 release after 10 min irradiation, see Section S4 of Supporting Information. Second, LA-AuNP slides were functionalized with an amino-ATTO550 dye such that the dye was covalently attached to the AuNP (ATTO550@AuNP) (Figures 2A and 3A and Supporting Information Figure S7). Imaging of this sample in the TIRF system, showed only photobleaching of the ATTO550 dye, presumably by interaction with the excited AuNP; in this case, the proximity is such that electron transfer (i.e., exchange) processes may contribute to degradation. Thus, the increments in emission are fully consistent with a release

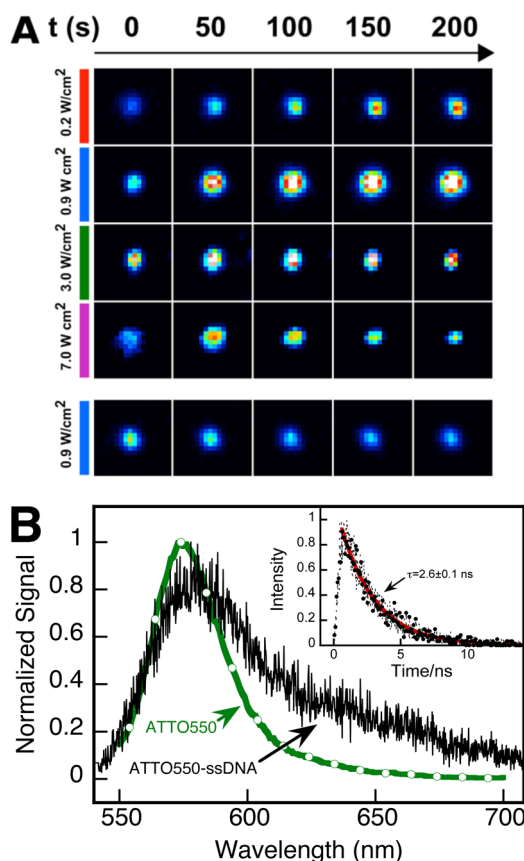


Figure 2. Real-time thermoplasmonic release of ssDNA from AuNP monitored in situ by TIRF microscopy using a 543 nm CW laser. (A) Representative fluorescence intensity images of ssDNA-ATTO550 release as a function of the irradiation time and the irradiance. The area of each image is $9 \mu\text{m}^2$ (pixel size 156 nm). The image uses a LUT false color scheme to represent the intensity of the fluorescence signal. The bottom sequence in panel A corresponds to a control using ATTO550@AuNP. (B) Emission spectra for ATTO550 free (green) and released from the dsDNA-AuNP conjugates (black) upon plasmon excitation. Inset: ATTO550 fluorescence decay under the same experimental conditions ($\tau_f \sim 2.6$ ns, data fit is shown in red).

mechanism due to specifically plasmon-induced melting of the noncovalent dsDNA interaction.

It is interesting to compare the intensity of the fluorescence spots in Figure 2A with the intensity curves in Figure 3A, bearing in mind that Figure 2A shows only representative spots, rather than average or ensemble data. This is particularly relevant for the series at 3.0 and 7.0 W/cm^2 . In both cases, the second image (at 50 s) is the more intense one, but at 7.0 W/cm^2 the loss of intensity is faster. The second image in Figure 2A corresponds to 150 and 350 J/cm^2 for 3.0 and 7.0 W/cm^2 , respectively. The corresponding curves in Figure 3A show maxima in the 200–250 J/cm^2 region in agreement with the images in Figure 2A, where the last image for 3.0 and 7.0 W/cm^2 corresponds to 600 and 1400 J/cm^2 , respectively; thus, it is not surprising that the latter shows faster bleaching or degradation.

In situ thermoplasmonic dsDNA melting dynamics were explored analyzing 50–90 bright spots from three independent experiments¹⁹ measured at different 543 nm CW laser irradiances ranging from 0.2 to 7.0 W/cm^2 . Supporting Information Figure S8 shows that ssDNA release becomes faster as the laser irradiance increases, whereas Figure 3A shows

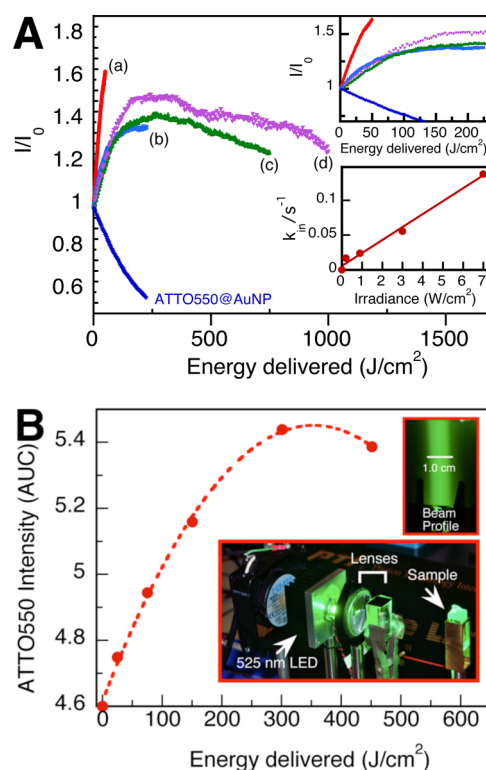


Figure 3. (A) Fluorescence intensity enhancement for ssDNA-ATTO550 release events as a function of total energy delivered. Average between 50 and 90 intensity vs time curves measured over single bright spots (see Figure 2) are plotted for each of the irradiances (a) red = 0.2, (b) blue = 0.9, (c) green = 3.0, and (d) purple = 7.0 W/cm^2 . ATTO550@AuNP conjugates were also tested as a control experiment (0.9 W/cm^2 , showing only emission decrease). Top inset: Expansion of the low energy region for Figure 3A. Bottom inset: Initial slope of the fluorescence intensity vs time for the experiments of Figure 3A as a function of the irradiance (curves shown in Supporting Information Figure S8). (B) ATTO550-ssDNA emission measured upon LED exposure of dsDNA@AuNP 1×1 cm quartz cuvettes (see section S4, Supporting Information). Top inset: shows the beam distribution of employed in our experimental setup (see bottom inset in this figure).

dependence with the total dose delivered. However, the top inset of Figure 3A shows that for irradiances at or above 0.9 W/cm^2 uniquely that only the total energy matters and not the form or rate of delivery. Only at very low irradiances (curve a in Figure 3A) is there a deviation from this behavior, as seen in the bottom inset of Figure 3A, with very low intensities being slightly more effective. This suggests a minor abundance (<2%) of an easily released population that is activated at low energies²⁰ but is too small to affect the overall observations at higher irradiances, where specific release due to cleavage of the noncovalent dsDNA interaction dominates. Further, 525 nm LED (22 mW/cm^2) in situ release experiments were carried out in 1×1 cm fluorescence cuvettes whose walls were modified with AuNP and the dsDNA, see Section S4, Supporting Information. These experiments showed similar behavior in terms of ATTO550 fluorescence intensity vs delivered power and those obtained at the single particle level, Figure 3B, with nondetectable Cy5 emission.

The melting temperature (T_m) of the dsDNA used in this study was 52 ± 2 °C at 10 mM NaCl; interestingly, this value increases to 66 ± 1 °C when the same dsDNA is bound to the

AuNP surface, which suggests an increment in the DNA packing onto the nanoparticle surface. Such of an increment in T_m is consistent with earlier studies¹³ (Figure 4). Clearly, upon

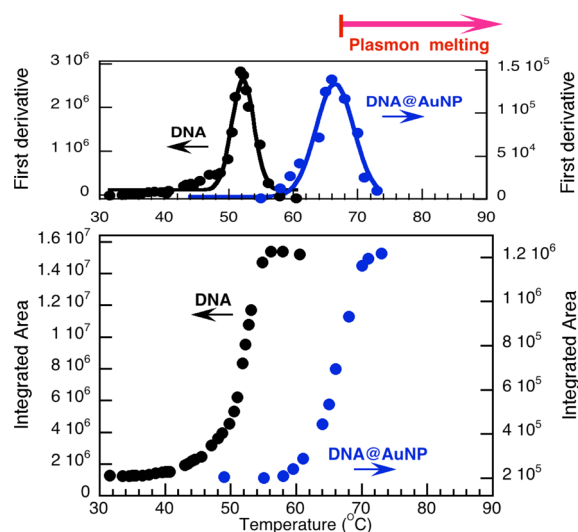


Figure 4. Melting curves for dsDNA in solution and when bound to AuNP using the strategy of Scheme 2. The top panel shows first derivative plots. The data correspond to the fluorescence increase as the temperature is raised. Note that the intensity values are smaller for DNA@AuNP (right curve in the plot), a measure of the low amount of DNA available at the surface and the difficulties of this measurement.

plasmonic excitation, the dsDNA bound to the AuNP shows effects similar to those observed at bulk temperatures that exceed 66 °C. The fact that the slopes of release curves at irradiances $>0.9 \text{ W/cm}^2$ are essentially identical, which suggests that the release mechanism does not change and is fully compatible with the ssDNA release discussed above. Thus, each photon arrival to a particle (approximately every 100 ns at $\sim 1 \text{ W/cm}^2$) has a constant probability of causing ssDNA release before the energy dissipates. Baffou and Quidant^{1,21} estimated that a 20 nm AuNP in water will experience a steady state 5 °C increase in temperature when exposed to $1 \text{ mW}/\mu\text{m}^2$ at 530 nm. Our irradiances are much lower than this, and the steady state temperature change should be well below 1 °C, thus insufficient to cause dsDNA dissociation. Thus, ssDNA release is not caused by bulk heating of the solution, but rather, it is a highly efficient surface-local event.²² It is possible that release events are favored at interparticle locations (hot plasmonic spots), as occurs in the case of plasmon-induced polymerization.^{23,24}

Further measurements at increasing salt concentrations did not show any statistical significance even at 1.2 M NaCl salt concentration.^{25,26} It seems likely that for successful release events, the temperature sensed by DNA is such that moderate changes in melting temperature do not change the release efficiency; instead, it seems that the packing of the dsDNA molecules onto the gold surface directs the melting dynamics of the oligonucleotide.

In summary, we report the in situ dynamics of thermoplasmonic ssDNA release from AuNP surface using fluorescence microscopy. Our findings point to the suitability of LA as a superior anchor able to support plasmon heating for ssDNA release.¹⁰ Our data indicate that the thermoplasmonic rate of ssDNA release is tunable by varying the excitation

irradiance, and that within the range examined, release is determined by the total energy delivered and not by the rate of delivery; quite significantly, this implies that although laser sources are convenient tools for our research, practical applications of this technology could be performed with simpler (as well as less expensive and safer) light sources, such as LEDs. Further, linearity with light source CW power is an asset for potential applications in sensors, therapeutics, or potential room-temperature miniaturized PCR devices. It is exciting to think about temporal and spatial control of PCR, as these are common characteristics for photochemical processes. Would having the materials attached to a nanoparticle enable cell-specific PCR? If the nanoparticle provides protection and a method for crossing cell membranes strategies could be developed to release photochemically siRNA in biological systems; a recent report suggests that this is indeed possible.¹² The demonstration of photocontrolled thermoplasmonic melting of DNA suggests that those possibilities are within reach. Clearly, there is room for optimization in relation to DNA length, tether length, and nanoparticle size and morphology; we were pleased to receive referee encouragement for us to follow some of these directions, something we hope to undertake in the future.

■ ASSOCIATED CONTENT

§ Supporting Information

Experimental details, characterization of AuNP on slides, DNA melting details, and effect of power on acquired images. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

The authors declare no competing financial interest.

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(20) We speculate that the 2% of relatively easy release may be due to a slight stoichiometric mismatch, that is, a slight excess of the ATTO-labeled chain. This chain could not hybridize due to the slight deficit of the complementary chain. It could still have some affinity for

the gold surface but be readily released upon thermoplasmonic heating well below the melting temperature of the assembly of Scheme 2.

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