Second Harmonic Super-resolution Microscopy for Quantification of mRNA at Single Copy Sensitivity

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ABSTRACT Cell-specific information on the quantity and localization of key mRNAs at single copy sensitivity in single cells is critical for evaluating basic cellular process, disease risk, and efficacy of therapy. Quantification of overexpressed mRNAs beyond the diffraction limit is constrained by the optical property of the probes and microscopy techniques. In this report, nanosized barium titanium oxide (BaTiO3, BTO) crystals were utilized as probes for mRNA quantification by a second harmonic super-resolution microscopy (SHaSM). The SHaSM was able to detect a single copy of the human epidermal growth factor receptor 2 (Her2) mRNA at a resolution of 55.6 nm with the ability to resolve multiple mRNA copies in a diffraction-limited spot. Her2 mRNA per cell was counted in SK-BR-3, MCF-7, and HeLa cell lines as 595 ± 79.1, 38.9 ± 8.26, and 1.5 ± 2.8, respectively. Our single-cell quantification results were validated with the fluorescence in situ hybridization studies and quantitative PCR, showing better specificity and selectivity over current single-molecule approaches for transcript detection. The SHaSM is expected to have an upper limit of resolving ∼10^4 transcripts in a single cell with the ability to monitor intracellular transcriptional dynamics at video rate. The developed approach has strong potential in clinical research and in the early diagnosis of life-threatening diseases such as cancer.

KEYWORDS: super-resolution microscopy · second harmonic generation · BaTiO3 nanocrystals · single-copy detection · mRNA quantification · diagnostics

Far-field light microscopy is the most desirable approach to “visualize” the microscopic world inside a cell or live organisms for biological studies. The ubiquitous property of noninvasive detection has made it extraordinarily versatile for investigating fine structural details of the cellular architecture. In addition, the use of a fluorophore further enables the study of subcellular dynamics ranging from molecular interactions to intracellular events.

Despite the revolutionary significance, the conventional far-field light microscopy is restricted by the diffraction limit based on the scale diffraction theory,¹ that is, objects within a diffraction-limited spot, ~250 nm, cannot be resolved. However, many subcellular structures have features much smaller than this limitation, and numerous intracellular dynamics occur within this space.² Several approaches have been developed to break the diffraction barrier to achieve super-resolution. One such effort is to localize individual fluorescent probes to obtain subdiffraction resolution.³–⁵ This stochastic super-resolution technique features two typical properties: the stochastic emission from single fluorescent molecules and the temporal modulation of the emission.⁶ These two features can be realized by either repeated photoactivation or photoblinking of fluorescent molecules, such as fluorescent dyes, proteins, or quantum dots. However, these fluorescent probes can easily undergo photobleaching under tens or hundreds of activation cycles.⁷ Quantum yield and photostability restrain the accuracy and applicability of this super-resolution technique.⁷ In addition, the blinking of the quantum dots limits the application of imaging subcellular dynamics.⁶ Furthermore, one-photon excitation configuration and total internal reflection imaging mode used in the stochastic super-resolution approaches constrain its application in deep-tissue imaging. These limitations demand the development of photostable probes and novel techniques for super-resolution imaging.

In this report, we demonstrate a novel second harmonic super-resolution microscopy...
(SHaSM), which was constructed on a scanning confocal microscopy platform taking advantage of ultrastable second harmonic generation (SHG) signals from nanosized crystals. Unlike the current super-resolution approaches which are based on fluorescent emission, the SHaSM collects ultrastable SHG emission from individual nanocrystals under the excitation of a two-photon laser source, overcoming the drawbacks of optical properties (photobleaching, photobleaching, quantum yield) and penetration depth of the fluorescence-based super-resolution imaging techniques.

SHG is a nonlinear optical process which converts two incident photons at a frequency \(ω\) into one photon at frequency \(2ω\). This process only occurs in materials which lack inversion symmetry. Under the illumination of an ultrafast laser, the SHG material emits a transient, high-intensity, nonblinking, non-illumination of an ultrafast laser, the SHG material in materials which lack inversion symmetry. Under the excitation of a two-photon laser source, overcoming the drawbacks of optical properties (photobleaching, photobleaching, quantum yield) and penetration depth of the fluorescence-based super-resolution imaging techniques.

**RESULTS AND DISCUSSION**

The SHaSM is a stochastic super-resolution technique and shares two typical features with the localization-based super-resolution microscopes: (i) sufficient photons are generated from single particles and (ii) stochastic modulation of the emission is possible. Figure 1a shows the SHG emission from individual BTO NCs deposited on a coverslip, where more than 3000 photons could be collected within a dwell time of 0.6 ms at each pixel. More importantly, the SHG emission from single BTO NCs is polarization-dependent. SHG emission of the BTO NCs stems from the distortion of oxygen atoms in the tetragonal unit cell (Supporting Information (SI) section 1). Since the BTO crystal has the symmetry of class 4, the SHG emission is only determined by the \(c\)-axis of the unit cell (Figure S1). Therefore, one can control the SHG emission by varying the angle between an incident electric field and the \(c\)-axis of the unit cell; meanwhile, polarization response analysis of the SHG signal can report on the orientation of the unit cell, which is beyond the scope of our study. Experimentally, we modulated the SHG signal from the BTO NCs by tuning the linear polarization angle of the excitation beam in the \(ij\) plane via a half-waveplate (SI section 1). Details of the polarization response of SHG NCs can be found in SI section 1 and Figure S1. Figure 1b shows the SHG emission from an individual BTO NC, marked as “1” in Figure 1a, with respect to the polarization angle.

The concept of the SHaSM is depicted in Figure 1c–f. Assuming that a ring consists of 12 BTO NCs shown in Figure 1f, single-molecule imaging of each BTO NC was realized via a conventional second harmonic imaging microscopy (columns 1 and 2 in Figure 1d). Temporal images are achieved by controlling the excitation polarization via a half-waveplate (column 2 in Figure 1d). Since the BTO NCs were stochastically dispersed, the second-order nonlinear susceptibility tensors of the BTO NCs randomly orient in the 3D space. Therefore, BTO NCs randomly emit a SHG signal with high efficiency under the excitation of a linearly polarized light. For each SHIM image, a point spread function (PSF) (eq 1) was used to localize the coordinates of the BTO NC.

In our work, we use BTO NCs as reporters for mRNA detection. Our strategy utilizes the concept of dimers formed from the hybridization of two BTO NC-conjugated probes targeting a mRNA in a sequence-specific manner. The dimer representing one single copy of mRNA will be resolved and quantified by the SHaSM with the spatial resolution of 5.5.6 nm, where it was possible to resolve and detect more than one mRNA in a single-diffraction-limited spot. This method can be applied to rigorously quantify the expression levels and the subcellular localization patterns of mRNAs at the sensitivity of a single copy, paving the pathway for quantitative monitoring of genetic information.
\[ P(x, y) = I_0 + A \exp \left[ \frac{1}{2} \left( \left( \frac{x - x_0}{s_x} \right)^2 + \left( \frac{y - y_0}{s_y} \right)^2 \right) \right] \]  

where \( P(x, y) \) is the 2D PSF on the \( xy \) plane, \( I_0 \) is the background intensity, \( A \) is the amplitude of the spot, \( x_0 \) and \( y_0 \) are coordinates of localization, and \( s_x \) and \( s_y \) are width of the PSF in \( x \) and \( y \) directions, respectively. By fitting, the coordinates can be localized with the standard deviation smaller than 1 nm (column 4 in Figure 1d). Super-resolution is therefore realized by superimposing all of the fitted coordinates of each BTO NC into one image (Figure 1e,f). Details of the algorithm can be found in SI section 2. The BTO ring, which cannot be resolved by scanning microscopy (Figure 1c), is identified by the SHaSM as shown in Figure 1f. Figure 1g shows the basic layout of the SHaSM instrumentation.

Figure 2 shows the proof-of-concept experiment of the SHaSM, which is validated by visualizing the BTO NCs beyond the diffraction limit. BTO NCs shown in Figure 1b and Figure 2b were processed by the SHaSM and led to a super-resolution image, as depicted in Figure 2e. For better comparison, BTO NC marked as spot “1” was elaborated with SHIM and SHaSM images shown in Figure 2a,d, respectively. In addition, the BTO NC marked as spot “2” was zoomed in, as shown in Figure 2c,f. Different from spot “1”, spot “2” has a much more complicated emission profile with respect to the incident polarization (SI Figure S2). This indicates that more than one BTO NC exists in the spot, which is confirmed by the SHaSM. Only a single BTO NC was modulated via a half-waveplate (10RPS52-2, Newport) mounted on a motorized rotation stage (PRM1Z8, Thorlabs) before being delivered onto the sample by a water-immersion objective (60x/1.20, Olympus). The SHG signal was collected using the same objective and further filtered by a short-pass filter and a band-pass filter (460-50, Chroma) before reaching the single-photon avalanche photodiodes (SPAD) (SPCM-AQR-14, PerkinElmer Inc.).
noted in spot ”1” (Figure 2d), while two BTO NCs were observed in spot “2” (Figure 2f). To characterize the SHaSM, the cross-sectional distributions of SHIM and SHaSM images are plotted in Figure 2g,h for spot “1” and “2”, respectively. The full width at half-maximum (fwhm) of spots in Figure 2a,c is ∼250 nm, the spatial resolution of the SHIM. In contrast, fwhm values of spots in SHaSM images are around 28 nm, about 9 times smaller than the diffraction limit. In addition, in Figure 2h, we show that the SHaSM can identify two adjacent BTO NCs with a center-to-center distance of 55.6 nm, which is the closest distance between two adjacent BTO NCs, as the first demonstration of second harmonic super-resolution imaging. Here we define the fwhm of a single spot (28.3 nm) as the localization precision of the SHaSM and 55.6 nm as the spatial resolution of the SHaSM. It should be noted here that in the SHaSM we localized the centroid of the 80 nm crystal, with the localization accuracy of ∼28 nm. This value is smaller than the size of the nanocrystal and allows us to distinguish two adjunct particles with the size of ∼80 nm. The estimated localization precision stems from the large-diameter BTO NCs since the SHG signal is evaluated based on the assembly of all of the unit cells in a single particle. This localization resolution can be improved to a few nanometers if smaller NCs are used, making the SHaSM a competitive super-resolution imaging approach with similar resolution as other super-localization/resolution microscopy techniques based on fluorophores or quantum dots.3–5,37

Given the evidence that the SHaSM can resolve single BTO NCs with a lateral resolution of ∼28 nm, we developed oligonucleotide probes with BTO NCs as ”reporters” for the detection of single mRNA beyond the diffraction limit. BTO NCs were initially chemically modified following the protocol in Figure 3a. Then thiol-terminated oligonucleotides were attached onto the surface-modified BTO NCs to form a BTO ”monomer” (Materials and Methods). We proposed a ”dimer” configuration of BTO NCs for mRNA detection instead of linearly hybridizing BTO ”monomers” due to the detection specificity. Two different oligonucleotide probes (probe strand (PS) and capture strand (CS)), which possess adjacent complementary sequences to the targeting mRNA, are conjugated to the BTO NCs. The 3′-thiolated end of PS and 5′-thiolated end of CS are connected to the surface of BTO NCs to form the two types of BTO ”monomers” (BTO_PS and BTO_CS). Finally, a BTO dimer can be formed upon hybridization of the mRNA (Figure 3b,d). In SHaSM, the dimer can be...
easily and visually seen from the images and differentiated from the background of individual NCs.

Before intracellular mRNA detection via the SHaSM approach, we prevalidated the hybridization of monomers to oligonucleotide sequences corresponding to a complementary mRNA sequence by in vitro experiments. A single-strand DNA (target strand, TS typically ~40 bps, depicting a target mRNA or miRNA) is targeted by a complementary PS (~20 bps) bearing a BTO particle and a CS (~20 bps) bearing a BTO particle consecutively to form a dimer upon hybridization. To illustrate the concept, BTO/C0 PS and BTO/C0 CS monomers were mixed with the solution of the TS (1:1:0.5) to result in a dimer configuration (BTO dimer) (Materials and Methods). Figure 3c,d shows scanning electron microscopy (SEM) images of BTO monomers and dimers spin-coated on the coverslip; individually distributed BTO monomers and dimers could be observed.

The in vitro calibration was performed to validate the SHaSM in differentiating BTO dimers from BTO monomers and for quantification. Mixture of BTO monomers and dimers in a 3:1 ratio were evaluated by SHaSM and further by SEM as the control. Figure 4a,b shows the SHIM image and SHaSM image of the mixture, respectively. The criterion for selecting a BTO dimer was based on the center-to-center distance between two monomers bearing the PS and CS hybridized to TS with a predefined interparticle spacing. Details of the algorithm used to detect and quantify BTO dimers are described in SI section 4. Distributions of BTO monomers and dimers separated from Figure 4b are shown in Figure 4c,d, respectively, and the zoomed SHIM and SHaSM images of BTO monomers and dimers are shown in Figure 4e–h. Results of the distance calculation between two monomers in over 50 BTO dimers are plotted in Figure 4i,j. Our analysis shows that the BTO dimers have an optimal interparticle distance of around 80 nm with an average at 77.36 ± 2.25 nm. The broad range of the interparticle distance of BTO dimers originates from two aspects; one is the size distribution of the BTO NCs, and the other is the geometry of the target strand (or mRNA).38 As a control, the dimer distance measured by SEM has a similar distribution as measured by SHaSM (Figure 4i); t test shows excellent agreement (p value is 0.31). In addition, BTO dimers were also quantified by SHaSM. Figure 4k compares the measured percentage of BTO dimers in a mixture of monomer and dimer by SHaSM (left) and SEM (right). In Figure 4k, left tilted is for the mixture with a dimer/monomer ratio of 3:1 and right tilted is for a ratio of 1:0. Our study shows that the results from two different evaluations compare well and are in good agreement. It should also be noted that in the in vitro quantification, the percentage of measured BTO dimers did not agree well with the designed ratio; this is because, in the in vitro hybridization, the yield of BTO dimers was not 100% and excess BTO monomers were not isolated from dimers. However, for detection in cells, the complementary oligonucleotide probes are expected to bind to the targeting mRNA and the nonspecific BTO monomers are washed away. Overall, Figure 4i–k successfully suggests that the SHaSM can be applied to detect and quantify BTO dimers with high accuracy.
Motivated by the capability of SHaSM to detect single BTO dimer dimers in vitro, we utilized our concepts and methodology to quantify mRNA corresponding to the human epidermal growth factor receptor 2 (Her2) at the single-cell level. It is known that the overexpression of the Her2 gene is associated with the tumorigenesis of 15–20% of invasive breast cancers and can be targeted by antibody-based therapies (e.g., Herceptin), whereas Her2-negative tumors require other therapeutic alternatives. The expression levels of Her2 mRNA have been used as predictive markers for diagnosis or in therapeutics for malignant tumors. Our proposed approach will provide quantitative information on the transcripts at single-copy sensitivity in single cells. For the intracellular mRNA hybridization, briefly the two types of BTO monomers (BTO_CS, BTO_PS) when incubated with a target (in vitro or in fixed single cells) will hybridize to form a dimer configuration; details of the experimental procedure are provided in the Materials and Methods section. The selection of cell lines is based on the estimated expression level of Her2 mRNA; it is been reported that Her2 mRNA is overexpressed in SK-BR-3 cells, moderately expressed in MCF-7 cells, and rarely expressed in HeLa cells. These cell lines were used as model systems to test the technology.

Figure 5 shows the expression of Her2 mRNA in SK-BR-3 cells (Figure 5a–c), MCF-7 cells (Figure 5d–f), and HeLa cells (Figure 5g–i). Panels a, d, and g in Figure 5 are the two-photon autofluorescence images that use 780 nm for excitation and 570–650 nm for emission; panels b, e, and h are SHG channel, and c, f, and i are merged images from both channels. For the two-photon and SHG microscopy, the laser power was set at 0.21 mW, resulting in an average intensity of 49.5 KW/cm² as well as a peak intensity of 5.96 GW/cm² at the focus, which is within the safety threshold for biological imaging. Although a minimum nonspecific binding was noted, our protocols show that all the free BTO NCs (or monomers) are washed away. In the following sections, we detail the quantification of Her2 mRNA in SK-BR-3 cells as an example to evaluate the distribution of mRNAs and the capabilities of the SHaSM.

Figure 5j shows the zoomed SHIM image of BTO NCs in SK-BR-3 cells marked in Figure 5c, and the corresponding SHaSM image is shown in Figure 5k. The algorithm for dimer differentiation was employed to quantify BTO dimers. Briefly, all BTO NCs were initially...
identified as red dots in SI Figure S8; later, any two closest BTO NCs with the center-to-center distance smaller than the threshold were selected as BTO dimers, which are noted as yellow dots in Figure 5l. A BTO dimer represents a single Her2 mRNA. From the magnified image of a single bright spot in the SHIM image (Figure 5m), three mRNAs can be identified from the SHaSM image, shown as red circles highlighted in Figure 5n. However, from the SHIM image, only one mRNA was noted. Here we should point out that the aggregate of multiple mRNAs is not the polycrystalline aggregate of BTO NCs, whose polarization responses are significantly different from that of single NCs. In our experiments, several treatment steps were performed to avoid polycrystalline aggregates. First, the surface of BTO NCs was chemically modified to ensure monodispersibility of NCs in solution, minimizing the possibility of aggregation. Second, the chemical groups (−NH2, −SH) on the surface of BTO NCs further prevent the formation of polycrystalline structures possibly due to aggregation. Last, the spherical shape of BTO NCs was chemically modified to ensure monodispersibility of NCs in solution, minimizing the possibility of aggregation.

Expression levels of Her2 mRNA were counted by the number of BTO dimers. SHaSM imaging of the whole cell can provide the quantity of transcripts, as shown in Table 1. The average number of Her2 mRNA in SK-BR-3 cells measured by SHaSM was estimated to be 595 ± 79.1 per cell. As a comparison, the expression measured by the conventional microscopy was 282 ± 54.3 per cell, which is more than two times lower than that quantified by SHaSM. The discrepancy exists because SHaSM has the ability to resolve multiple mRNAs in a diffraction-limited spot which otherwise is not possible by conventional microscopy. The magnified SHaSM images demonstrate that it is possible to locate more than one mRNA in a diffraction-limited spot should these exist.

Quantification of Her2 mRNA in MCF-7 cells and HeLa cells was also performed via the SHaSM, and the results are provided in Table 1. Our analysis shows that the average number of Her2 mRNA quantified by the SHaSM is 38.9 ± 8.26 per cell in MCF-7 and 1.5 ± 2.8 per cell in HeLa cells, which agrees well with the past values. However, from the SHIM images, the value noted was 153.1 ± 31.9 per cell in MCF-7 and 3.4 ± 5.8 per cell in HeLa. We are of the opinion that the discrepancy mainly stems from the background of nonspecific BTO monomers, which may bind to the active sites in the cell cytoplasm. Similar defects have also been reported for the FISH technique. The SHaSM has the capability to exclude the interference from nonspecifically bound BTO monomers, and consequently, unless the Her2 mRNA is targeted by two BTO monomers, to result in a dimer formation, the
monomers will not be counted. From Table 1, it is our opinion that, in the conventional diffraction-limited mRNA quantification approach, the mRNA number is overestimated in cell lines with moderate or rare expression due to the nonspecifically bound probes, while the mRNA number is underestimated in overexpressing cell lines because of the inability to resolve several targets in one spot.

To validate our SHG probes and detection strategy, FISH experiments were performed along with conventional qPCR to compare the expression level of Her2 mRNA in the above cell lines. Results from qPCR show that the expression level of Her2 mRNA in the SK-BR-3 cell line is much higher than that observed in MCF-7 and HeLa cell lines (SI Figure S9); in addition, immunostaining for the Her2 protein on the cell membrane also indirectly suggests that Her2 mRNA is over-expressed in SK-BR-3 cells (SI Figure S11). Furthermore, quantitative FISH images, as shown in SI Figure S10 and Table 1, indicate that the average number of Her2 mRNA in SK-BR-3, MCF-7, and HeLa cell lines is comparable to the results detected by the SHaSM and is close to the theoretical estimates reported by other works via FISH.41,42

Overall, the SHaSM approach offers significant accuracy and specificity compared to the conventional diffraction-limited detection methods for transcript quantification in single cells, especially for cells with highly expressed mRNA transcripts. The theoretical maximum number of SHG probes detected by the SHaSM is about 8–10 in each diffraction-limited spot (≈250 nm in diameter), resulting in a probing range of approximately 40–50 μm², suggesting the theoretical quantification limit of 10,000–15,000 transcripts per cell. Furthermore, this quantification limit can be extended with a polarization analyzer (typically inserted before the detector) to improve the contrast between different polarization angles. By SHaSM, we will have the ability to “see” individual BTO NCs and thus “visualize” single mRNA, which is almost impossible by other super-resolution microscopy techniques due to the optical constraints posed by fluorescent probes. It is worth noting that in our design the SHG probes have minimal requirements on the length of the targeting mRNA and hence are suitable for the detection of any single mRNA with a short sequence length; in addition, only two BTO NCs (BTO_CS and BTO_PS) are required to target each transcript to detect and quantify Her2 mRNA; this design is also possible to implement via plasmonic nanoparticles43 and hyper-spectral super-resolution approach.44,45

The localization precision and lateral resolution of SHaSM in our experiment was estimated to be ~28 and ~56 nm, but in theory, these two values can reach single digits. In a typical scanning image with a 200 μs dwell time, N = ~2000 photons can be collected from a single BTO NC to yield the theoretical spatial resolution as 1/√N.46 However, in our experiment, the lateral resolution is restricted due to the volume of the BTO NCs. In addition, the SHaSM also has the potential for rapid imaging. By optimizing the photons collected in each pixel, we can improve the scanning speed of the SHIM image to 500 frames per second, but still with high SNR ratio (>100). Even if every 50th SHIM image is processed to obtain a super-resolution map by the second harmonic generation process, at least 10 frames of the super-resolution SHG image can be realized per second, which is sufficient to monitor the dynamics of intricate intracellular behavior compared to nonfluorescent approaches.43,47–49 Furthermore, we anticipate that the imaging speed can reach video rates if scan-free instrumentation is used in the future.14

CONCLUSIONS

To summarize, we present a SHaSM platform by taking advantage of SHG signals from BTO NCs to demonstrate single-molecule sensitivity at a resolution of a few tens of nanometers. In vitro experiments consistently demonstrate the capability of SHaSM to differentiate two individual BTO NCs separated by a center-to-center distance of 55.6 nm with a localization resolution of ~28 nm. The BTO NCs were functionalized with complementary oligonucleotide sequences to hybridize with target-specific mRNA for visualization at super-resolution scale and quantification at single-copy sensitivity. The SHaSM approach conceptualized was utilized to not only visualize the localization patterns but also to quantify the expression levels of individual Her2 mRNA in fixed SK-BR-3, MCF-7, and HeLa cell lines, suggesting that the number of copies of Her2 mRNA in the respective cell lines were 595 ± 79.1, 38.9 ± 8.26, and 1.5 ± 2.8, respectively, and was in good agreement with the theoretical calculations and validated with FISH and qPCR experiments. We expect the quantification technology will allow scientists to “see” individual mRNA transcripts inside a single cell and to elucidate gene expression patterns in specific phenotypes at different stages of tissue differentiation. Such knowledge will be paramount for human health as defects in transcription adversely affect normal growth and differentiation, which will impact both clinical diagnostics and basic biology.

MATERIALS AND METHODS

Chemicals and Reagents. Thiol-modified and simple DNA oligonucleotides were obtained from IDT (Coralville, IA). The oligonucleotides were of highest quality, purified either using HPLC or from PAGE gels. Succinimidyl-4-(N-maleimidomethyl)-cyclohexane-1-carboxylate (SMCC) was purchased from Pierce.

residual surfaces were treated with ethanol. Next, the solution was washed with nanopure water. Oligosequences were prepared and stored at 4 °C. After successful conjugation, all prepared probe-strand-conjugated BTO NCs were washed three times with 0.3 M PBS to remove unbound oligonucleotides. After the unconjugated BTO NCs were removed by centrifugation, the DNA NCs were mixed with 2 mM of SMCC dissolved in DMSO and kept at room temperature for 2 h. Free SMCC was removed by centrifugation.

**BTO-Oligonucleotide Hybridization.** Three different oligonucleotide strands (TS, PS, and CS) were designed using NCBI Blast, and these oligonucleotides were purchased from IDT-DNA, Inc. (Coralville, IA). The 3'-thiolated end (probe strand) and 5'-thiolated end (capture strand) oligonucleotides were used for the conjugation with BTO NCs; each oligonucleotide is made up of 10 base pairs of linkers along with 20 base pairs of complementary sequences. In addition, a complementary oligonucleotide target strand for the probe strand and capture strand was also used here to form an in vitro dimer structure as a control. A probe strand (PS): 5'-AGAAGCTGAGTAGGGTTGGGTTTTTTTTTTTTCCTCACC-3', and a capture strand (CS): /ThiolM/-5'C0O-C176-tttttttttttGTTCCTCTACCC-CTAAGTGAC-3'.

We adopted the protocol of gold nanoparticle hybridization with oligonucleotides previously used in our group with minor modifications.3,5,11 DNA oligonucleotides were first dissolved in 10:1 TE (Tris-EDTA) buffer to make a 100 μM solution; the disulfide bond of thiolated DNA oligonucleotide probe was then broken using a 10 mM solution of TCEP (at room temperature, shaking for 10–15 min). These reduced thiolated DNA strands (1 μM) were then reacted with 1 mL of the functionalized (thiol-reactive) BTO NC solution to obtain 1 mL of DNA-BTO conjugate solution (at ~60 °C for 30 min to 1 h). The residual surfaces were finally blocked with 10 mM mercaptoethanol. Next, the solution was washed with nanopure water (18.2 MΩ, Milli-Q, Inc.). Finally, the resulting DNA–BTO probes were washed three times with 0.3 M PBS to remove unbound oligonucleotides. After successful conjugation, all prepared probes were stored at 4 °C until further use.

For control experiments, BTO dimers in vitro, constituting probe-strand-conjugated BTO NCs and capture-strand-conjugated BTO NCs, were mixed in equal proportions, and then the target strand was added in lower concentration than the conjugated BTO NCs (2:1); the mixture was kept at a temperature of 60 °C, which was lower than the melting point of the DNA hybridization. Electron microscopy (Figure 3) and dynamic light scattering (SI section 1) experiments were used to characterize the function and conjugation of BTO probes.

**Mature Probe and Incubation with BTO NCs.** The standard culture media used for SK-BR-3, MCF-7, and HeLa was Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum. Cells were seeded onto the sterilized no. 1 coverslips (VWR International, Batavia, IL) inside a 6-well plate with culture medium and maintained at 37 °C in an atmosphere of 5% CO2; after reaching 80% confluence, cells were rinsed with PBS three times and then fixed by 4% paraformaldehyde for 15 min; after being washed by PBS three times, cells were permeabilized by 0.1% Triton-X for 2 h; in the end, cells were washed by PBS.

For mRNA detection, fixed cells were incubated with functionalyzed BTO–oligonucleotide probes (probe strand and capture strand at the same concentration) at room temperature overnight in the PBS and then washed with gentle shaking and stored in PBS at 4 °C for SHG imaging.

**Conflict of Interest:** The authors declare no competing financial interest.

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**Supporting Information Available:** Additional information and figures. This material is available free of charge via the Internet at http://pubs.acs.org.
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