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Single-nucleobase resolution of a surface energy transfer nanoruler for *in situ* measurement of aptamer binding at the receptor subunit level in living cells†

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In situ identification of aptamer-binding targets on living cell membrane surfaces is of considerable interest, but a major challenge, specifically, when advancing recognition to the level of membrane receptor subunits. Here we propose a novel nanometal surface energy transfer (NSET) based nanoruler with a single-nucleobase resolution (SN-nanoruler), in which FAM-labeled aptamers and single-sized gold nanoparticle (GNP) antibody conjugates act as a donor and an acceptor. A single nucleobase resolution of the SN-nanoruler was experimentally illustrated by molecular size, orientation, quenching nature, and other dye-GNP pairs. The SN-nanoruler provides high reproducibility and precision for measuring molecule distance on living cell membranes at the nanometer level owing to only the use of single-sized antibody-capped GNPs. *In situ* identification of the aptamer binding site was advanced to the protein subunit level on the living cell membrane for the utilization of this SN-nanoruler. The results suggest that the proposed strategy is a solid step towards the wider application of optical-based rulers to observe the molecular structural configuration and dynamic transitions on the membrane surface of living cells.

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Introduction

Cell systematic evolution of ligands by exponential enrichment (cell-SELEX) is an exciting method that enables the creation of cell-specific aptamers without prior identification of a specific target and expands the various applications of aptamers in diagnostics, drug delivery, therapeutics, *etc.*¹ Precise identification of desired aptamer binding sites is more important and still considered the main challenging scientific issue, the

solution of which will greatly promote further aptamer development in the future. To address this issue, various analytical methods are conducted. For example, chromatography² and mass spectroscopy^{3,4} have evidenced that aptamer Sgc8 optimally binds to membrane protein tyrosine kinase 7,⁵ AS1411 recognizes nucleoli proteins,^{6,7} SYL3C recognizes the epithelial cell adhesion molecule,^{8,9} XQ-P3 binds to programmed death ligand 1, *etc.* Additionally, XQ-2d, a pancreatic ductal adenocarcinoma PL45 cell-binding aptamer, has been confirmed to bind the transferrin receptor protein (CD71).^{10,11} Interestingly, the aforementioned methods are complicated, tedious, labor-intensive, costly, and all performed *in vitro* and failed to accurately determine the specific protein subunit at the aptamer recognition site due to the limitations of the measuring principle. Thus, the measurement of aptamer binding sites on the surface of intact living cells *in situ* is necessary for understanding the exact biological functions of aptamers. There is an urgent need to develop *in situ* strategies for living cell systems, particularly when advancing recognition to the protein subunit level.

Optical-based distance measurements at the nanoscale, referred to as “nanoruler”, are one of the key topics in cell biology, promising a potential tool for *in situ* determination of ligand-binding sites. The most commonly used nanoruler is Förster resonance energy transfer (FRET), as proposed by Förster *et al.* in 1948, but it is limited to a scale of less than 10 nm.

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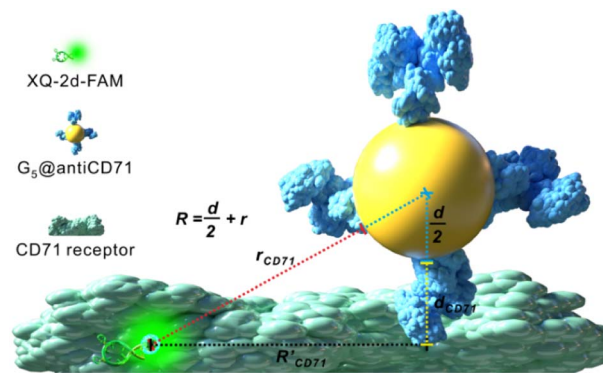
Quenching efficiency (Φ) of FRET follows the law $\Phi = 1/[1 + (r/r_0)^6]$.¹² Alternatively, nanometal surface energy transfer (NSET) is an energy transfer process between a nanometal surface and fluorescent dyes, as proposed by T. L. Jennings *et al.* in 2005. The Φ of NSET is speculated to follow the law $\Phi = 1/[1 + (r/r_0)^4]$.¹³ Similar to FRET, Φ is the energy transfer efficiency, r_0 is the distance at which a specific dye displays equal probabilities for energy transfer and spontaneous emission, and r is the separation distance from the fluorophore to the surface of GNPs or from the donor dye to the receptor dye.^{14,15} The angstrom resolution of FRET has been shown by rigorously considering molecular distances and orientations at the single-molecule level.¹⁶ In principle, NSET can measure much longer distances than FRET¹³ and could become an essential nanoscale analytical tool.¹⁷ For instance, NSET has been innovatively designed to detect the conformational changes in RNA binding¹⁸ and to explore the interaction between DNA and fluorescent dyes,^{19,20} but the resolution limits of its distance measurements have not been addressed experimentally. Therefore, developing new models of the NSET nanoruler with a known-resolution is crucially needed to expand its potential for *in situ* accurate measurement on living cells, especially for the scalability and possibility of measuring more refined spatial configuration.

We herein propose a novel *in situ* NSET-based nanoruler with single-nucleobase resolution (hereinafter shortened to SN-nanoruler), composed of an FAM-labeled aptamer and single-sized GNP-antibody conjugates as the donor and acceptor, respectively, on the living cell membrane surface. Taking into account molecular distance, orientation, quenching nature, *etc.*, single-nucleobase resolution of the SN-nanoruler is evidenced by both fluorescence intensity (FLI) and lifetime (FLT) experiments. By inserting an “n TA” base spacer between the dye and 5' end of the XQ-2d aptamer, the Φ values gradually increase by approximately 6% for lengthening of each per base. Application of single-sized antibody-capped GNPs enables ultra-high reproducibility and accuracy during distance measurements. Size of antibodies here is considered by three methods: theoretical calculations, TEM and AFM measurements, respectively. The error coming from antibody measurements by the three methods causes a non-significant difference on further distance measurements, which evidences the feasibility of theoretical calculations on antibody size. High single-nucleobase resolution of the SN-nanoruler enables advancing aptamer binding site *in situ* identification to the protein subunit level on living cell membranes.

Results

Design principle of the SN-nanoruler

The principle of the SN-nanoruler can be illustrated by selecting an XQ-2d aptamer and its target CD71 receptor as a model system on the PL45 cell membrane surface (Scheme 1). Although GNPs with a larger size could accept energy more efficiently, GNPs with a smaller size were reported to be the least intrusive for living cells.¹⁴ Also, GNPs with a diameter of 5 nm (G_5 NPs) have simplicity and reproducibility in both chemical synthesis and surface modification procedures.^{21,22} Importantly,



Scheme 1 SN-nanoruler working model for measuring distance on the CD71 receptor of PL45 cells. R represents the distance from FAM dye to the center of G_5 NPs, r represents the distance from FAM dye to the surface of G_5 NPs, and R' represents the separation distance between the XQ-2d-FAM binding site and antiCD71 binding site. The d and d_{CD71} are the diameters of G_5 NPs and antiCD71 antibody, respectively.

the r_0 has already been determined to be 13 nm for the G_5 NP and FAM dye pair with a high agreement between theoretical predictions and experimental data.¹⁵ Thus, XQ-2d labeled with FAM dye (XQ-2d-FAM) acts as an energy transfer donor, while single-sized G_5 NPs act as energy transfer acceptors. Notably, FAM is far from an ideal fluorophore in the living cell system; its emission might be influenced by surrounding factors. Nevertheless, more information on the NSET of GNP-FAM pairs could be referred to in many previous studies; more importantly, the emission of FAM completely overlaps with the absorption of G_5 NPs which is an important parameter in the NSET model.^{15,23,24} Hence, here FAM was selected for constructing the SN-nanoruler. As evidence, a very stable but expensive dye, Alexa488, was labelled on the 5' end of XQ-2d and showed similar quenching behaviors in the SN-nanoruler model although its r_0 value is unknown to calculate the exact distance, which will be discussed later.

Accordingly, G_5 NPs were functionalized with saturated antiCD71 on their surface ($G_5@antiCD71$), and the 5' end of XQ-2d was labeled with the FAM fluorophore. After XQ-2d and antiCD71 respectively carried the FAM fluorophore and G_5 NPs to their binding sites on the CD71 receptor protein, the energy transfer was monitored by flow cytometric analysis on FLI and time-correlated single photon counting measurement on FLT, respectively. A single nucleobase resolution of the SN-nanoruler was experimentally illustrated by molecular size, orientation, quenching nature, and other dye-GNP pairs. Here, antibody size (d_{Ab}) was estimated by TEM, AFM and theoretical prediction which will be discussed in detail later, and was taken into account by assuming a spatial configuration of a right triangle.

$$\left(\frac{d}{2} + d_{Ab}\right)^2 + R'^2 = R^2 \quad (1)$$

$$R = r + \frac{d}{2} \quad (2)$$



Importantly, the SN-nanoruler working model is the interaction process of a point dipole and surface dipole,²⁵ where the donor (dye) is viewed as the point dipole and the acceptor (GNP) models as a collection of infinite point dipoles,²⁶ and consequently, the equation of separation distance-dependent quenching efficiency proposed by Yun *et al.* is as follows:¹³

$$\Phi = \frac{1}{1 + \left(\frac{r}{r_0}\right)^4} \quad (3)$$

Eqn (3) exhibits the relationship between Φ and r . In particular, for the FAM-G₅NP pair, Φ could be calculated as r_0 has been determined to be 13 nm,¹⁵ which means that these parameters can be acquired according to the change in the FLI of XQ-2d-FAM before and after the binding of G₅@antiCD71.

The anisotropy of ligand binding is an important parameter affecting the NSET process. A C₆ chain is introduced as a spacer between the FAM dye and 5' end of each aptamer (Table S1†). This flexible spacer offers a cone of probability for randomizing the dipole vector relative to the nanometal surface, assures an isotropic distribution of electronic dipoles, and guarantees that the measurement result is not affected by dye orientation.¹⁴

FLI-based quenching could be dynamic quenching or static quenching. So, it is necessary to find a way to examine the nature of fluorescence quenching. Here, the Stern–Volmer quenching constant (K_{sv}) was examined that enjoy the relationship as follows:²⁷

$$\frac{F_0}{F} = 1 + K_{sv}[Q] \quad (4)$$

where $[Q]$ is the amount of the quencher, F_0 is FLI without a quenching agent, and F is FLI after adding quenchers. For

dynamic quenching, F_0/F and $[Q]$ follow a good linear relationship, while for static quenching, this relationship is a rising curve bent to the y-axis. Furthermore, the FLT of the donor dye was examined before and after the presence of the quencher. For dynamic quenching, the relationship between F_0/F and τ_0/τ follows

$$\frac{F_0}{F} = \frac{\tau_0}{\tau} \quad (5)$$

where τ_0 is the FLT without a quenching agent and τ is the FLT after adding quenchers. Here's an advance note: the NSET process of the SN-nanoruler is dynamic quenching rather than static quenching, evidenced by both the FLI and FLT experiments, which will be discussed in detail later.

Single-nucleobase spatial resolution of the SN-nanoruler on living cell membranes

Firstly, we examined the binding features of two ligands on PL45 cells. PL45 cells were uncontaminated with high integrity of the cell nucleus as evidenced by confocal imaging (Fig. S1†). PL45 cells were treated with a series of concentrations of XQ-2d-FAM, examined by flow cytometry (FCM) (Fig. 1A and B). The result indicated that the binding saturation concentration of XQ-2d-FAM is 150 nM. Notably, although the used concentration of XQ-2d-FAM is saturated, which is much less than the self-quenching concentration,^{28,29} it effectively avoids the self-quenching that happened. Then, the optimal incubation time for XQ-2d-FAM was verified to be 30 min through incubating 150 nM XQ-2d-FAM with these cells for different times (Fig. 1C and D). And the calculated equilibrium dissociation constant (K_d) is of ~11 nM, consistent with our previous report, indicating that the labeling of the XQ-2d aptamer by the FAM fluorophore does not affect its high binding affinity on the PL45

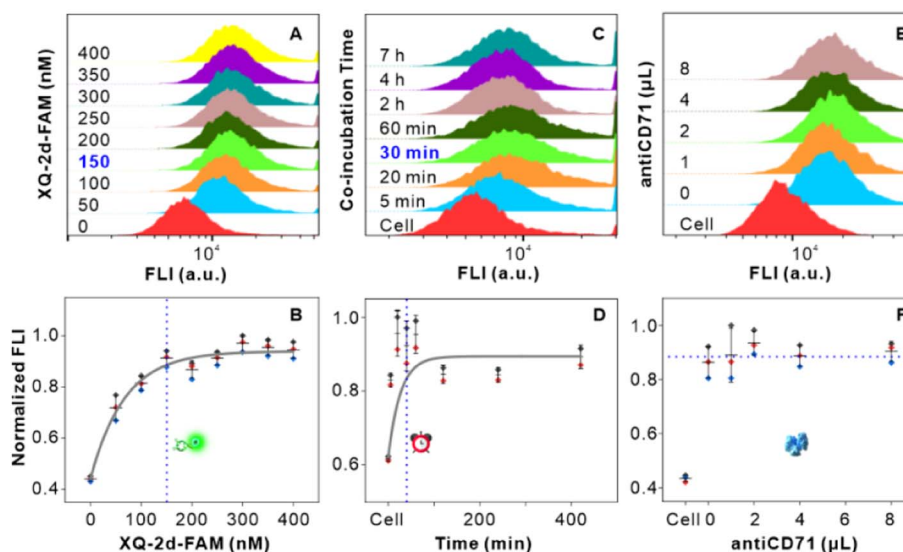


Fig. 1 Binding feature analysis of the ligands in living PL45 cells. (A and B) Saturated binding concentration analysis of the XQ-2d-FAM aptamer to the CD71 receptor on PL45 cells. The FCM charts indicated 0, 50, 100, 150, 200, 250, 300, 350, and 400 nM of XQ-2d-FAM from bottom to top in A, respectively. (C and D) Best co-incubation time of the XQ-2d-FAM aptamer with the CD71 receptor on PL45 cells. The FCM charts indicated native PL45 cells and cells with saturated XQ-2d-FAM for about 5, 20, 40, and 60 min and 2, 4, 7 and h from bottom to top in C, respectively. (E and F) Competitive binding analysis of saturated XQ-2d-FAM and different amounts of antiCD71, including 1, 2, 4, and 8 μ L, respectively.



cell membrane (Fig. S2†). Also, competitive binding between the aptamer XQ-2d-FAM and the monoclonal antibody antiCD71 was further excluded by FCM (Fig. 1E and F). Increasing amounts of antiCD71 were incubated with PL45 cells for CD71 receptor protein recognition to compete with saturated XQ-2d-FAM, but no decrease in the FLI of FAM was observed, demonstrating non-competition between XQ-2d-FAM and antiCD71. Moreover, another competitive test was carried out by incubating PL45 cells first with different amounts of antiCD71 and then with saturated XQ-2d-FAM (Fig. S3A and B†). It also evidenced that XQ-2d-FAM and antiCD71 binding to two distinct sites of the CD71 receptor protein, *i.e.*, the CD71 receptor, antiCD71 and XQ-2d-FAM could act as ideal components on the living cell membrane to examine the validity of the SN-nanoruler.

Secondly, we precisely conjugated G₅NPs and antiCD71. Antibody is amphoteric, pH impacts the antibody adsorbed onto GNPs.³⁰ Thus, we explored the pH effect on conjugating

G₅NPs with antiCD71, which evidenced a good stability and high conjugation efficiency at pH 7.5 (Fig. 2A). Then, the saturated amounts of antiCD71 were incubated with 200 μ L of G₅NPs to maximize the conjugation of antibody on the GNP surface (Fig. 2B), which indicated that 1.2 μ L of antiCD71 could reach saturation binding in 200 μ L of the G₅NP system. The successful conjugation of G₅NPs with antiCD71 was evidenced by UV-VIS absorbance and agarose gel electrophoresis (Fig. 2C), as well as zeta potential analysis (Fig. S4†). Compared with native G₅NPs, G₅@antiCD71 had a larger diameter of hydration dynamics and the poly-dispersity index (PDI) of DLS assays increased from 0.1 to 0.3 after conjugation (Fig. 2D); nevertheless, the results demonstrated a good dispersibility of G₅@antiCD71. GNPs, antibody, and aptamer are the key components in the calculation of separation distance. In the SN-nanoruler, sizes of G₅NPs and antibody were considered when calculating the distance between the energy donor and receptor. TEM, AFM, and theoretical calculations were

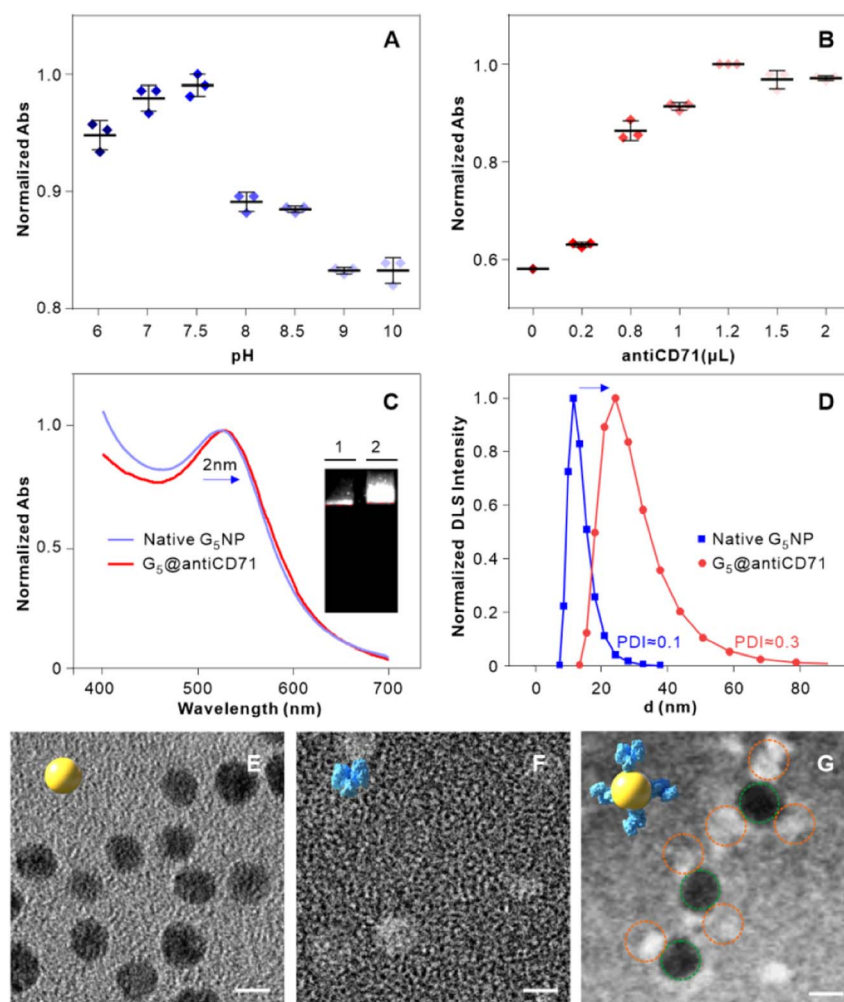


Fig. 2 Preparation and characterization of a G₅@antiCD71 conjugate. (A) Normalized absorbance of the G₅@antiCD71 conjugate at varied pH values. (B) Normalized absorbance of the G₅@antiCD71 conjugate for various volumes of antiCD71. (C) UV-VIS absorption spectra of native G₅NPs (blue line) and G₅@antiCD71 (red line), respectively. The inset indicates agarose gel electrophoresis assay of native G₅NPs (channel 1) and G₅@antiCD71 (channel 2). (D) The polydispersity index on hydration dynamics diameters of native G₅NPs and G₅@antiCD71 by DLS measurements, respectively. (E–G) TEM images of native G₅NPs, native antiCD71, and G₅@antiCD71, respectively. The green and yellow circles in G indicate G₅NPs and antiCD71, respectively. The scale bar in (E–G) is 5 nm.



respectively used to estimate the sizes of antiCD71 and G_5 -@antiCD71. Negative-staining TEM analysis evidenced that the diameters of G_5 NPs, antiCD71, and G_5 @antiCD71 were 5.5 ± 0.4 nm, 5.2 ± 0.2 nm, and 10.0 ± 1.0 nm, respectively (Fig. 2E–G and S5†). Notably, although the coating of antiCD71 has heterogeneity, the protein corona does not affect the properties of GNPs, such as dispersion and stability.³¹ Therefore, antiCD71 antibody protein corona coating on GNPs can carry the GNPs to the site on the CD71 receptor but does not affect the reliability and accuracy of this SN-nanoruler function. AFM observed that the sizes of G_5 NP, antiCD71 and G_5 @antiCD71 were 5.3 ± 0.6 nm, 4.9 ± 0.6 nm, and 9.0 ± 1.2 nm, respectively (Fig. S6A–D†). Notably, Stokes has proposed a theoretical calculation of protein sizes according to eqn (6) (ref. 32).

$$R_{\text{Stokes}} = 66 \times 10^{-3} M^{\frac{1}{3}} \quad (6)$$

where R_{Stokes} is the Stokes radius and M is the atomic mass unit of the protein. Consequently, the size of antiCD71 antibody with an M of 95 KD is *ca.* 6.0 nm. Statistical analysis showed no significant differences in the sizes of antiCD71 determined by TEM, AFM, and the theoretical calculation (Fig. S7†). The theoretical calculation effectively supported the reliability of the experimental results, and its deviation affecting final separation distance calculations will be discussed later.

Thirdly, we measured the specificity and validity of the SN-nanoruler at the CD71 receptor protein level in living PL45

cells. We verified the incubation time of G_5 @antiCD71 coupling with PL45 cells. FCM analysis showed that the FLI of PL45 cells gradually dropped and reached a plateau after co-incubating with G_5 @antiCD71 for 3 h (Fig. S8†), which was chosen as the co-incubation time of G_5 @antiCD71 in the following assays. Moreover, confocal imaging visualized the NSET effect of XQ-2d-FAM on G_5 @antiCD71. XQ-2d-FAM clearly bound to the membrane of PL45 cells (Fig. 3A and B). The staining of the cell nucleus with 4,6-diamidino-2-phenyl indole (DAPI) dye further evidenced the membrane surface binding of XQ-2d-FAM (Fig. 3C and D). Next, a significant fluorescence quenching was observed after adding G_5 @antiCD71 (Fig. 3E), compared to a negative control of G_5 @BSA conjugates (Fig. 3F), which verified the high specific quenching from G_5 @antiCD71. Furthermore, the FLI of XQ-2d-FAM before and after the binding of G_5 @antiCD71 was quantitatively analyzed using grayscale maps (Fig. 3G–J), indicating a fluorescence quenching of about over 60%. Confocal imaging showed unavoidable large errors in FLI analysis but clearly evidenced the specificity and validity of the SN-nanoruler.

We further deduced the feasibility and specificity of the SN-nanoruler by FLI and FLT related experiments. Quantitated FLI was estimated by FCM. During the whole assay, PL45 cells had good monodispersities before and after the binding of XQ-2d-FAM and different amounts of G_5 @antiCD71, respectively (Fig. S9†), and the FCM charts are presented in Fig. 4A. The Φ could be written as

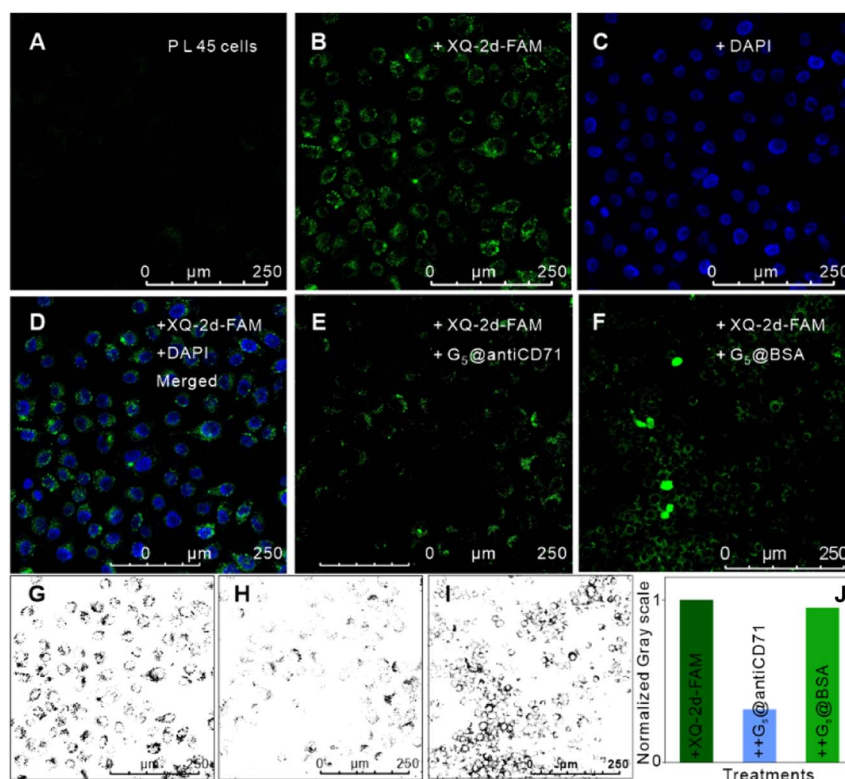


Fig. 3 Feasibility verification of the SN-nanoruler by confocal imaging analysis of PL45 cells. (A) Native PL45 cells without the fluorescence background. (B) PL45 cells with the saturated XQ-2d-FAM aptamer. (C) The cell samples in (B) further treated by DAPI staining. (D) The merged image of (B) and (C). (E) PL45 cells treated with excess saturated G_5 @antiCD71, after incubation with saturated XQ-2d-FAM. (F) PL45 cells treated with excess saturated G_5 @BSA, after incubation with saturated XQ-2d-FAM, as a negative control. (G–I) Grey values of cell FLI in (B), (E), and (F) that were estimated using Image J software. (J) Normalized grey values of (G–I) that were estimated using Image J software.



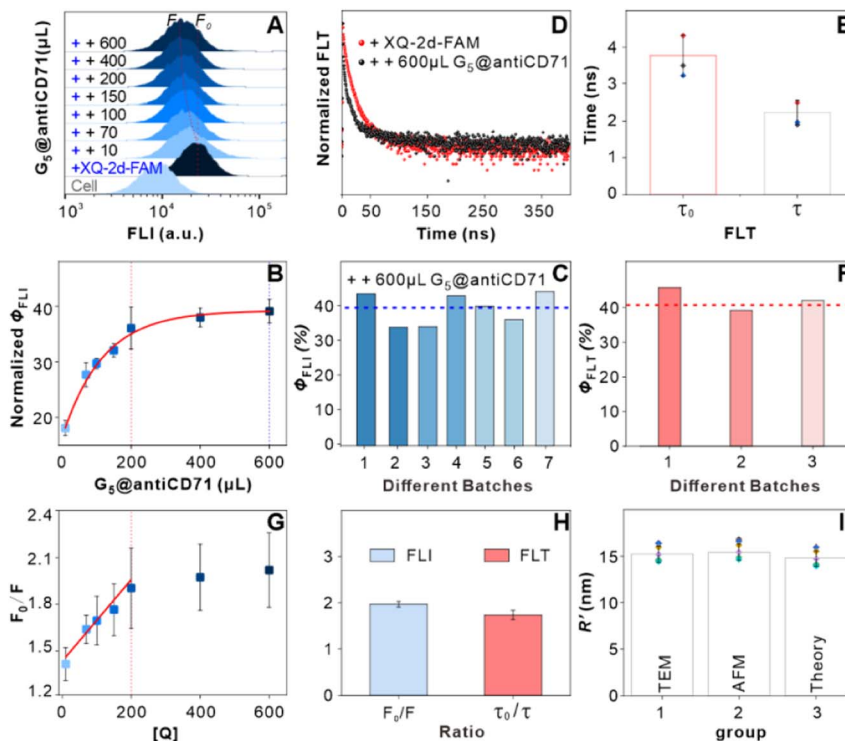


Fig. 4 Feasibility verification of the SN-nanoruler by FLI and FLT related experiments. (A) FCM analysis on PL45 cell samples. FCM charts indicated native PL45 cells (cell), cells with saturated XQ-2d-FAM (+FAM), and cells with saturated XQ-2d-FAM and $G_5@antiCD71$ of different amounts, *i.e.*, 10, 70, 100, 150, 200, 400, and 600 μL from bottom to top, respectively (from “+ 10” to “+ 600”). (B) Fluorescence quenching efficiency as a function of the amounts of $G_5@antiCD71$. (C) Seven repeats of the Φ test that were calculated by FLI experiments. (D) FLT of cells incubated with saturated XQ-2d-FAM (+FAM, red line) and then co-incubated with excess saturated $G_5@antiCD71$ (+ $G_5@antiCD71$, black line). (E) Mean FLT of cells with saturated XQ-2d-FAM treatments before (τ_0) and after co-incubation with the saturated $G_5@antiCD71$ quencher (τ), respectively. (F) Three repeats of the Φ test that were calculated by FLT experiments. (G) Linear relationship of F_0/F and $[Q]$ when added with different amounts of $G_5@antiCD71$. (H) The ratio of F_0/F calculated using FLI (blue) and τ_0/τ calculated using FLT (red), respectively. (I) Analysis of distance between two binding sites in the presence of saturated XQ-2d-FAM and saturated $G_5@antiCD71$ according to the estimated antibody sizes from TEM (group 1), AFM (group 2) and the theoretical calculation (group 3), respectively.

$$\Phi = \frac{I_0 - I}{I_0} \quad (7)$$

where I_0 is the FLI of XQ-2d-FAM bound to PL45 cells minus native PL45 cells and I is the FLI of XQ-2d-FAM in the presence of $G_5@antiCD71$ minus native PL45 cells. As displayed in Fig. 4B, the Φ reached a plateau when the amounts of $G_5@antiCD71$, labelled as $[Q]$, reached over 200 μL , and to guarantee the stability of quenching, 600 μL of $G_5@antiCD71$ was chosen for the following experiments. For more accurate estimation of quenching efficiency, the experiment was carried out in seven different batches (Fig. 4C), and the calculated Φ is $39 \pm 4.59\%$. Quantitated FLT was evaluated by time-correlated single photon counting measurements (Fig. 4D); the mean FLT of XQ-2d-FAM bound to PL45 cells was 3.77 ns (τ_0), and the subsequent saturation binding of $G_5@antiCD71$ made its lifetime decrease to 2.23 ns (τ), *i.e.*, the FLT decreased by $42 \pm 3.29\%$ (Fig. 4E and F). Interestingly, F_0/F follows a good linear relationship of $[Q]$ dependence, *i.e.*, $y = 0.00265x + 1.40952$, when the $[Q]$ is no more than 200 μL , and F_0/F reached a stable value of ~ 1.9 when $[Q]$ is larger than 200 μL (Fig. 4G and the blue bar in Fig. 4H). These results implied the nature of dynamic fluorescence quenching as discussed in the above section.

Moreover, the calculated mean τ_0/τ is ~ 1.8 (red bar in Fig. 4H), nearly equal to the ratio of F_0/F , which further evidenced dynamic quenching rather than static quenching during this NSET process. Obviously, any Φ value can be utilized to numerically calculate a R' ; however, only when fluorescence quenching is maximal and stable is the value of R' closer to true physical imaging. In an FLI related experiment, when Φ is 39%, the value of R'_{CD71} was 15.22 ± 0.78 nm, 15.43 ± 0.77 nm, and 14.81 ± 0.74 nm, respectively, based on the parameters determined by TEM, AFM, and theoretical calculation, and statistical analysis showed no significant difference among the three methods for determining R'_{CD71} (Fig. 4I), which were all similar to the previously reported.¹¹ Interestingly, the measurement error in R'_{CD71} by the SN-nanoruler was about 4.29% from TEM analysis, 4.75% from AFM analysis, and 4.53% from the theoretical calculation, all smaller than previously reported errors of at least 10%.¹¹ About 5% measurement error in R' can be considered acceptable random fluctuations,³³ so the SN-nanoruler is a relatively accurate spectral measurement tool in the field of chemo-metrics. Therefore, these results not only evidenced the specificity and validity of the SN-nanoruler, but also indicated that the antibody size could be theoretically



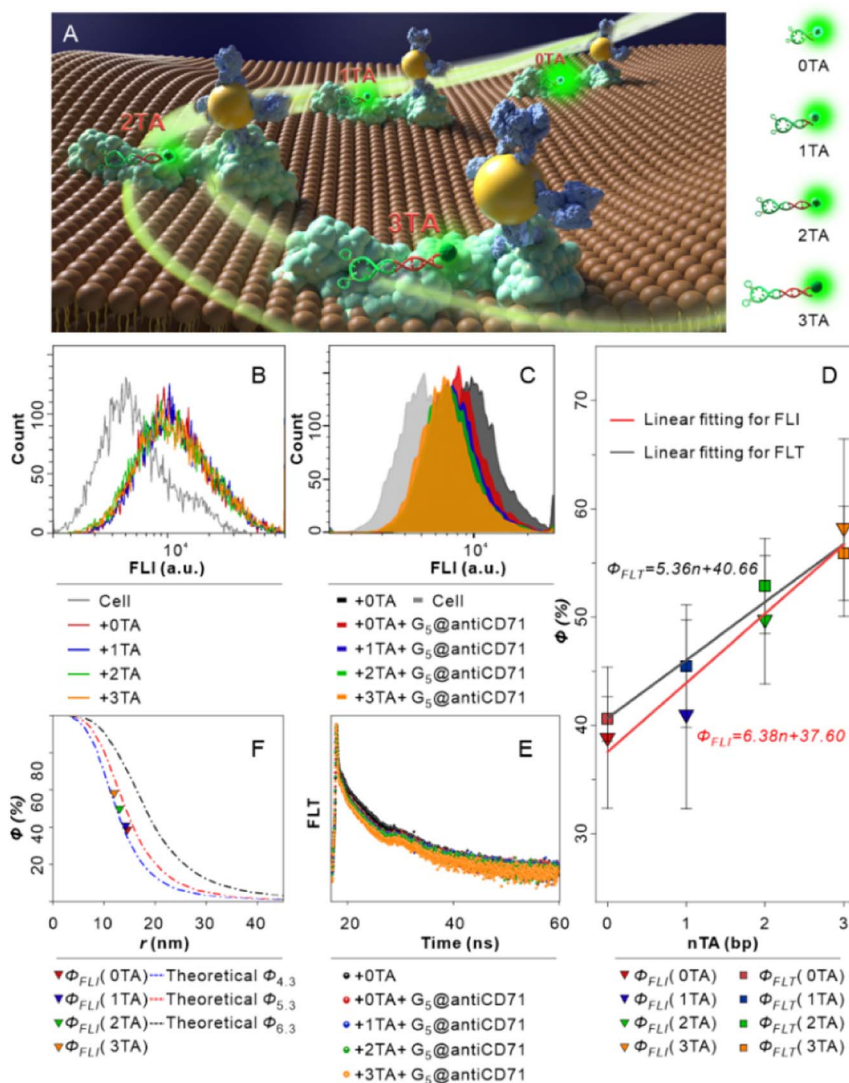


Fig. 5 SN-nanoruler with a single-nucleobase resolution identified by FLI and FLT experiments. (A) Single-nucleobase resolution model by lengthening the “ n TA” spacer, *i.e.*, one TA base pair at a time, between the 5′ end of XQ-2d and FAM, where n is the number of TA base pairs. (B) FCM analysis of native PL45 cells (cell), cells incubated with saturated XQ-2d-FAM (0 TA), XQ-2d-1TA-FAM (1 TA), XQ-2d-2TA-FAM (2 TA), and XQ-2d-3TA-FAM (3 TA), respectively. (C) FCM analysis of cells, cells incubated with saturated 0 TA, and cells incubated with saturated n TA and then saturated G₅@antiCD71, respectively. (D) Linear fitting relationship between the number of TA and Φ_{FLI} (red line) and Φ_{FLT} (black line), respectively, calculated according to the change in FAM FLI and FLT. (E) FLT analysis of cells incubated with saturated 0 TA and cells incubated with saturated n TA and then co-incubated with excess saturated G₅@antiCD71, respectively. (F) Comparisons of our experimental data with the theoretical simulation curves. Theoretical curves show the r -dependent Φ of GNPs with diameters of 4.3 nm (blue line), 5.3 nm (red line), and 6.6 nm (black line), respectively, adopted from the literature.⁴⁵ The color triangles indicate experimental Φ values calculated from FLI.

estimated without laborious TEM or AFM experiments in the future.

A nanoscale ruler with high spatial resolution is essential in materials engineering, genetics, biochemistry, and medicine,³⁴ while the spatial resolution of the NSET effect has not been experimentally explored before. Here the SN-nanoruler model enables its spatial resolution to be examined by lengthening the “ n TA” spacer between the 5′ end of XQ-2d and FAM dye, *i.e.*, XQ-2d- n TA-FAM where n is the number of TA bases (hereinafter shortened to 0 TA, 1 TA, 2 TA, and 3 TA, Fig. 5A). Importantly, the 0 TA, 1 TA, 2 TA, and 3 TA binding to PL45 cells generated similar fluorescence features (Fig. 5B), and their K_d values were all lower than 15 nM (Fig. S10A–F†), indicating a strong binding

affinity of 0 TA, 1 TA, 2 TA, and 3 TA to PL45 cells, and the slight difference did not affect the FLI under saturation binding. Previous reports also found that terminal modification did not alter the binding characteristics of the aptamer.^{35,36}

The change in FLI was monitored after insertion of 1, 2, and 3 TA bases. In detail, PL45 cells were incubated with saturated 0 TA, 1 TA, 2 TA, and 3 TA, subsequently co-incubated with G₅@antiCD71, washed, then re-suspended with binding buffer, and detected by FCM. The Φ_{FLI} of the 0 TA group is approximately 39%, the same as the results in the above section. Then, the Φ_{FLI} gradually increased to 41%, 50%, and 58% for 1 TA, 2 TA, and 3 TA groups, respectively (Fig. 5C). Clearly, the insertion of the “ n TA” spacer decreased the separation distance between



the FAM fluorophore and the metal surface, n TA-dependent Φ_{FLI} fitting showed a good linear relationship, *i.e.*, $\Phi_{\text{FLI}} = 6.38n + 37.60$, and the slope of linear fitting is 6.38% (red line in Fig. 5D), indicating that the Φ_{FLI} value increased by 6.38% when lengthening every TA base pair.

Similarly, the change in the FLT was further monitored after the insertion of 1, 2, and 3 TA base pairs (Fig. 5E). By co-incubation with the saturated $G_5@antiCD71$, the mean FLT of 0 TA, 1 TA, 2 TA, and 3 TA groups gradually dropped, from 3.77 to 2.23, 2.04, 1.76, and 1.65 ns, respectively (Fig. S11†). Correspondingly, the Φ_{FLT} gradually increased from 41% that is similar to the result in the above section, to 46%, 53%, and 56%, respectively, following an “ n TA”-dependent linear relationship, *i.e.*, $\Phi_{\text{FLT}} = 5.39n + 40.66$ (black line in Fig. 5D), which is very similar to the results of FLI. There is a slight difference in the FLI and FLT results which may originate from operation and/or different instrumentation.

Three theoretical r -dependent Φ curves of GNPs with diameters of 4.3 nm (blue line), 5.3 nm (red line), and 6.6 nm (black line) adopted from the literature¹⁵ were plotted and compared with the experimental data corresponding to different “ n TA” spacers (color triangle), and these data were overall located between the $G_{4.3}NP$ and $G_{5.3}NP$ theoretical curves (Fig. 5F), although Φ_{FLI} (0 TA) with a value of 39% has a slight larger deviation. Actually, NSET has the highest sensitivity when the Φ is around 50%. When hypothesizing that r is nearly the value of r_0 with a slight variation of Δr , Φ will be around 50%, and r can be written as

$$r = r_0 + \Delta r \quad (8)$$

According to eqn (3),

$$\Phi = \frac{1}{1 + \left(\frac{r_0 + \Delta r}{r_0}\right)^4} \quad (9)$$

If Δr is considered as the change of r when lengthening every TA base pair, the insertion of 0, 1, 2 and 3 TA base pairs exactly induced the changes of Φ around 50%, and the corresponding r is 14.56 nm, 14.24 nm, 13.03 nm, and 11.96 nm, respectively, with a mean Δr of ~ 0.87 nm. As reported, the pitch of the double DNA helix is 0.34 nm per base pair because of the double chain rigidity, while the extension of single-stranded DNA on the GNP surface is reported to be 0.15 nm per base because the orientation of ssDNA was more flexible compared to that of dsDNA.³⁷ The length per base for oligonucleotides was estimated to be about 0.43 nm,³⁸ while the calculated Δr is larger than the reported extension length per base pair.

Two factors may contribute to the above deviation: one factor may be the inaccurate estimation of previously reported r_0 , especially in the living cell system. Ignoring higher order terms, eqn (9) can be approximately written as

$$\Phi \approx \frac{1}{2 + 4\frac{\Delta r}{r_0}} \quad (10)$$

The reconfigured equation is as follows:

$$r_0 = \frac{4 - 8\Phi}{\Phi} \Delta r \quad (11)$$

Obviously, r_0 is directly proportional to Δr when Φ is a known value around 50%. Actually, the r_0 could be affected by many factors as proposed by C. J. Breshike *et al.*¹⁵

$$r_0 = \frac{\alpha\lambda}{n_m} (A\phi)^{\frac{1}{4}} \left(\frac{n_r}{2n_m} \left(1 + \frac{\varepsilon_1^2}{|\varepsilon_2|^2} \right) \right)^{\frac{1}{4}} \quad (12)$$

where α is the orientation of the donor to the GNP receptor, λ is the maximum emission wavelength of the donor, A is the absorptivity of a thin film mirror, ϕ is the quantum yield of the donor, and n_m is the refractive index of the solvent system, n_r is the refractive index of the GNP receptor, ε_2 is the complex dielectric function of the GNP receptor, and ε_1 is the solvent dielectric. In this work, optimized r_0 of the FAM- G_5NP pair that is reported in *in vitro* aqueous solution was applied; however, the different surface capping of GNPs might alter the refractive index of the system (n) and the dielectric function (ε), especially in the living cell system.

The other factor is the $\Phi(r)$ power law relationship. As discussed in the above section, when $\Phi = 1/[1 + (r/r_0)^4]$ is applied, the donor is seen as a point dipole while the GNP is seen as an infinite surface and a film mirror.¹⁴ C. J. Breshike *et al.* still thought that Φ and r exhibited a four-power law behaviour after redefining r_0 according to the complex dielectric function (ε) and absorptivity (A).¹⁵ In fact, the FAM-labeled aptamer is not a perfect point dipole, and the G_5NP acceptor is neither an infinite surface nor a perfect mirror, but a sphere with a curved surface and size only a few times larger than the dye. Thus, the four-power law of $\Phi(r)$ relationship may overestimate the interaction distance between the G_5NP acceptor and the FAM dye donor. Nevertheless, all of the above data evidenced the capability of the SN-nanoruler for distinguishing single-nucleobase resolution measurement, even under the conditions of the living cell environment and the fluidity of the cell membrane.

Just in case, other dye-GNP pairs were considered. Alexa488, a quite stable dye commonly used for *in situ* analysis, was further used as the energy donor to verify the resolution of the SN-nanoruler. As shown in Fig. S12A,† the insertion of the “ n TA” spacer did not alter the binding affinity of XQ-2d-Alexa488 to PL45 cells. After the saturated binding of XQ-2d- nTA -Alexa488, PL45 cells showed a similar FLI (Fig. S12B†). Subsequently, when co-incubated with saturated $G_5@antiCD71$, the insertion of the “ n TA” spacer caused the Φ_{FLI} to increase from 34% to 40%, 53%, and 55%, respectively (Fig. S12C and D†), following an “ n TA”-dependent linear relationship, *i.e.*, $\Phi_{\text{FLI}} = 7.56n + 33.82$. Obviously, the quenching feature curve of Alexa488 dye on G_5NPs is different from that of the FAM dye, *i.e.*, the Φ_{FLI} value increased approximately 7.56% when lengthening every TA base pair. So far, the r_0 value of Alexa488- G_5NP is unknown, but could be deduced from the n TA-dependent Φ values if assuming that the base-pair



lengthening exactly equals the distance separating. It may pave a new way to determine the unknown r_0 value of other dye-GNP pairs, which will be discussed in our on-going work. High precision, *i.e.*, single-nucleobase spatial resolution of this SN-nanoruler exhibited high potential for *in situ* identification of aptamer-binding sites and advanced the molecular distance measurement further to the level of protein subunit domains in the living cell system.

In situ measurement of aptamer binding sites at the protein subunit level in living cells

The above results of high single-nucleobase resolution inspired us to advance the *in situ* identification of aptamer binding sites to a much deeper level, *e.g.*, protein subunits, in living cells. Recently, our group selected new aptamer DML7 that binds to DU145 cells.³⁹ Preliminary data of SDS-PAGE, as well as pull-down and confocal analysis, hinted that its potential target might be integrin $\alpha 3\beta 1$,⁴⁰ a transmembrane heterodimer protein consisting of the α subunit domain and the β subunit domain.⁴¹ However, further distinguishing the protein subunit is very challenging with current technology. In this context, the integrin $\alpha 3\beta 1$ receptor protein, anti $\alpha 3$ (P1B5) antibody, anti $\beta 1$

(K-20) antibody, and DML7-FAM aptamer were used for conducting the SN-nanoruler to *in situ* measure the separation distance on the DU145 cell membrane (Fig. 6A). It may provide a clue to the probable binding subunit of integrin $\alpha 3\beta 1$. Notably, we first determined the saturation concentration of DML7-FAM binding to DU145 cells. DU145 cells were cultivated and shown to have good integrity of cell morphology and the nucleus by confocal microscopy (Fig. S13†). DU145 cells were then treated with different concentrations of DML7-FAM, followed by washing with washing buffer, suspending in binding buffer, and testing through FCM. The results indicated that the saturation concentration of DML7-FAM binding to DU145 cells is 100 nM (Fig. 6B and C). Similar to the above studies, we further determined the optimal time of DML7-FAM incubation with DU145 cells. DU145 cells were co-incubated with 100 nM DML7-FAM at different times. The saturation of FLI suggested that the optimal time for DML7-FAM binding with DU145 cells is 60 min (Fig. 6D and E).

An important foundation of the SN-nanoruler is the non-competitive binding between two ligands when observing the distance between their binding sites. For this purpose, competition assays between the DML7-FAM aptamer and anti $\alpha 3$

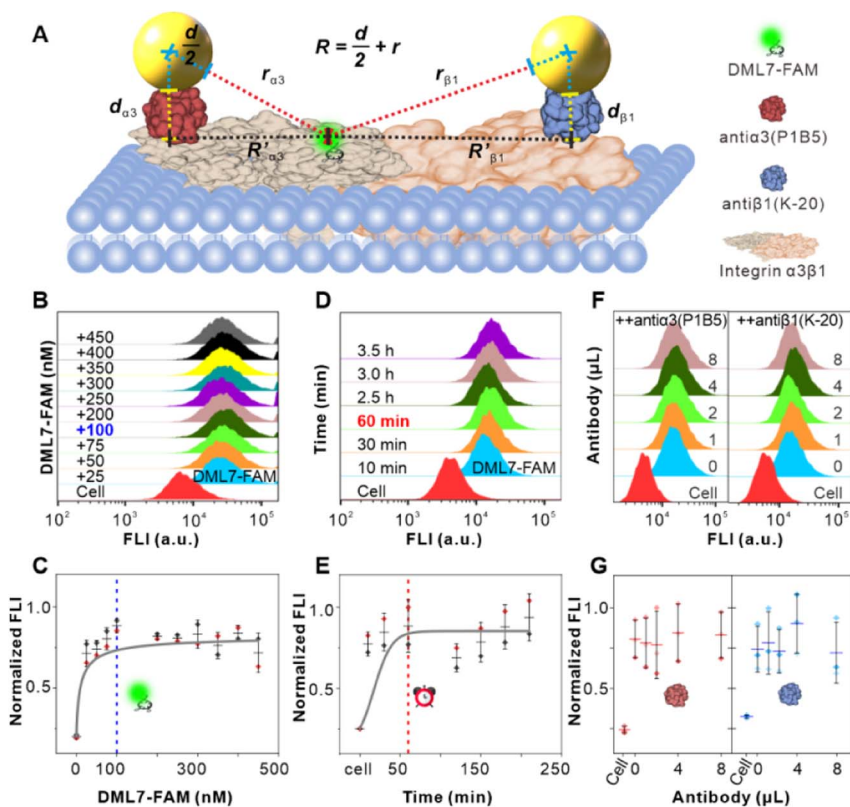


Fig. 6 Binding feature analysis of DML7-FAM and two antibodies on the integrin $\alpha 3\beta 1$ receptor of DU145 cells. (A) Illustration on the SN-nanoruler for measuring $\alpha 3\beta 1$ subunit-level distance. DML7 and each antibody bring FAM dye and G₅NP to their binding sites, respectively. $R'_{\alpha 3}$ and $R'_{\beta 1}$ represent the separation distance between the corresponding binding sites, and $d_{\alpha 3}$ and $d_{\beta 1}$ represent the size of each antibody, respectively. (B and C) Saturated binding concentration analysis of DML7-FAM. The numbers in the FCM chart indicated each final concentration (nM) of DML7-FAM incubated with native DU145 cells (cell). (D and E) Best co-incubation time analysis of DML7-FAM. The numbers in the FCM chart indicated each incubation time of saturated DML7-FAM with native DU145 cells (cell). (F and G) Competitive binding analysis of saturated DML7-FAM and different amounts of anti $\alpha 3$ (P1B5) (left) and anti $\beta 1$ (K-20) (right) including 1, 2, 4, and 8 μ L, respectively. And the order of incubation with cells is DML7-FAM and antibody.



(P1B5) or anti β 1 (K-20) antibody were respectively monitored by FCM. After saturation binding with the saturated DML7-FAM aptamer, DU145 cells were continuously incubated with various amounts of unlabeled anti α 3 (P1B5) antibodies. It showed no obvious decline of FLI (left panel of Fig. 6F and G), demonstrating no competition between DML7-FAM and anti α 3 (P1B5) antibody. Similarly, the results showed no competition between DML7-FAM and anti β 1 (K-20) antibody (right panel of Fig. 6F and G). To further exclude the possibility of co-recognition between DML7-FAM and anti α 3 (P1B5) or anti β 1 (K-20) antibodies, DU145 is first incubated with antibodies and then with the aptamer, *i.e.*, changing the incubation order of DML7-FAM and antibodies (Fig. S14A–D†). The incubation sequence did not affect the FLI of DML7-FAM bound to DU145 cells, indicating that DML7-FAM and the two antibodies have distinctly different binding sites on the DU145 cell membrane. Moreover, the competitive assay was examined by simultaneously adding DML7-FAM and each antibody to the cell system (Fig. S15A–D†). All these results indicated no competition between DML7-FAM and each antibody, further excluding nonspecific interference between different ligands. Similarly, the optimal pH value is near 7.0 for the coupling of G_5 @anti α 3 (P1B5) (Fig. 7A) and G_5 @anti β 1 (K-20) (Fig. 7B). The UV-VIS

absorbance spectroscopy, agarose gel electrophoresis (Fig. 7C and D), and zeta potential (Fig. S16A and B†) results evidenced the successful coupling of each G_5 @antibody. Furthermore, 0.2 μ L of anti α 3 (P1B5) and 0.4 μ L of anti β 1 (K-20) were determined to be excess in 200 μ L of G_5 NP solution to make a saturated conjugation (Fig. S17A and B†), and the DLS results demonstrated a good dispersibility of G_5 @anti α 3 (P1B5) and G_5 @anti β 1 (K-20), respectively (Fig. 7E and F).

In particular, negative-staining TEM images confirmed that native anti α 3 (P1B5), G_5 @anti α 3 (P1B5), native anti β 1 (K-20), and G_5 @anti β 1 (K-20) have sizes of 7.1 ± 0.8 nm, 11.9 ± 1.2 nm, 6.8 ± 1.1 nm, and 12.5 ± 0.2 nm (Fig. 7G–J), respectively, and AFM images evidenced their sizes of 6.9 ± 0.6 nm, 12.9 ± 1.9 nm, 6.6 ± 0.8 nm, and 13.0 ± 2.4 nm, respectively (Fig. S18A–F†). According to eqn (6), the theoretical sizes of anti α 3 (P1B5) and anti β 1 (K-20) with an M of 150 KD and 138 KD were 7.0 nm and 6.8 nm, respectively. Related sizes of native antibody and G_5 @antibody and the R' value are summarized in Table 1. Statistical analysis showed no obvious difference among the measurement results by these three measurement methods (Fig. S19A and B†).

Confocal imaging first visualized the variation of NSET efficiency between DML7-FAM and different G_5 NP-antibody

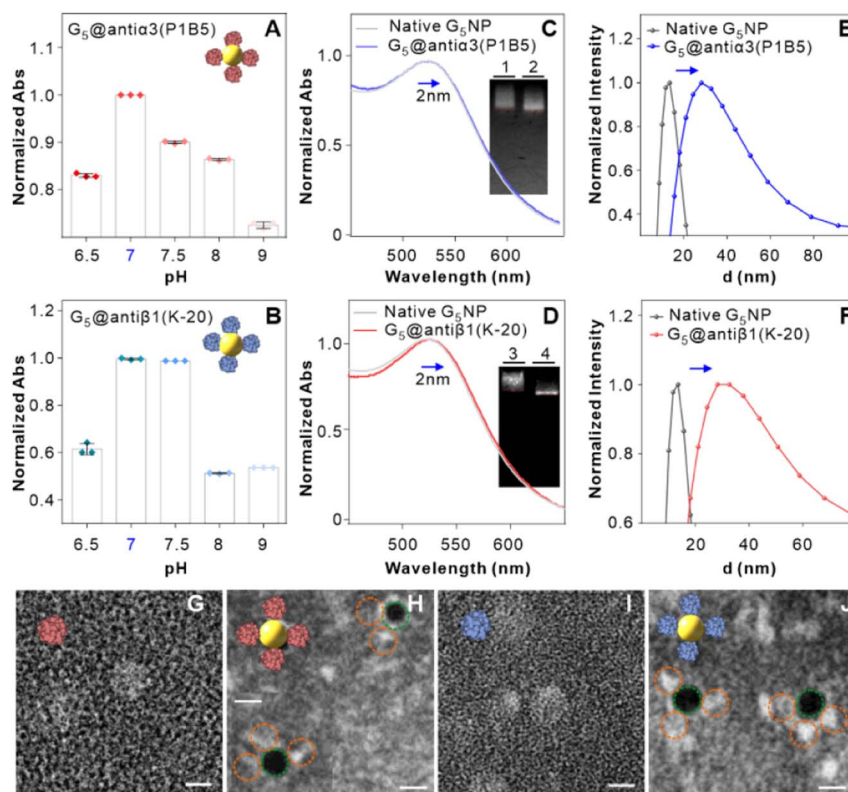


Fig. 7 Synthesis and characterization of the G_5 @antibody conjugate. (A and B) Normalized absorbance of anti α 3 (P1B5) and G_5 @anti β 1 (K-20) conjugates at varied pH values. (C and D) Normalized UV-VIS absorption spectra of native G_5 NP (gray line), G_5 @anti α 3 (P1B5) (blue line), and G_5 @anti β 1 (K-20) (red line). The insets of C and D indicate agarose gel electrophoresis assay of G_5 @anti α 3 (P1B5) and G_5 @anti β 1 (K-20) (channels 1 and 3) and native G_5 NPs (channels 2 and 4). (E and F) The polydispersity on hydration dynamics of native G_5 NPs (gray line), G_5 @anti α 3 (P1B5) (blue line) and G_5 @anti β 1 (K-20) (red line), respectively. (G–J) TEM images of native anti α 3 (P1B5), G_5 @anti α 3 (P1B5), native anti β 1 (K-20), and G_5 @anti β 1 (K-20), respectively. The green and yellow circles in (G–J) indicate G_5 NPs and antibodies, respectively. The scale bar in (G–J) is 5 nm.



Table 1 Sizes of G_5 @antibody and mean separation distance R'_{mean}

Name	anti α 3 (P1B5)	anti β 1 (K-20)	G_5 @anti- α 3 (P1B5)	G_5 @anti- β 1 (K-20)
TEM	7.1 ± 0.8	6.8 ± 1.1	11.2 ± 0.7	12.1 ± 1.5
AFM	6.9 ± 0.6	6.6 ± 0.8	12.9 ± 1.9	13.0 ± 2.4
Theory	7.0	6.8	12.3 ± 0.6	12.1 ± 0.6
Tukey HSD ^a	n.s	n.s	n.s	n.s
R'_{mean}	—	—	14.1	16.1

^a n.s. represents no significant difference.

conjugates. Fig. 8A–C illustrate the binding of DML7-FAM on the surface of DU145 cells, but a significant fluorescence quenching of DML7-FAM was observed when adding saturated

G_5 @anti α 3 (P1B5) (Fig. 8D) or G_5 @anti β 1 (K-20) (Fig. 8E). G_5 @BSA conjugates as the negative control did not alter the FLI of DML7-FAM on DU145 cells which clearly excluded the

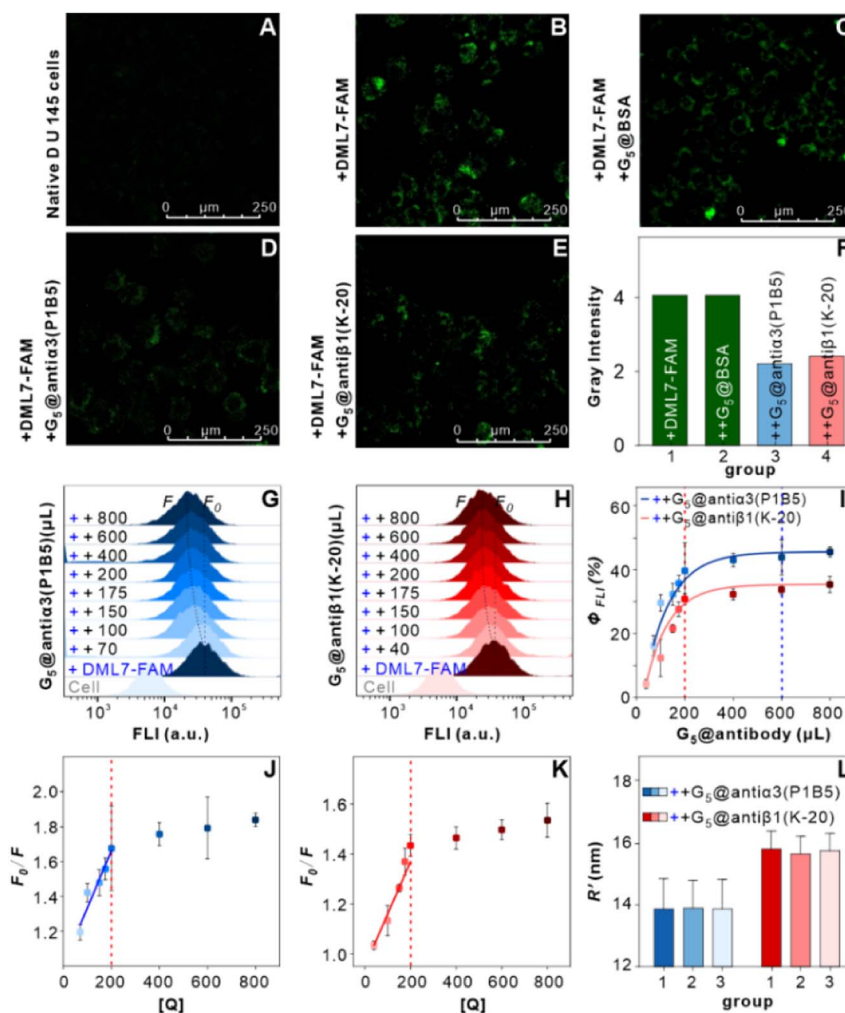


Fig. 8 Measurement of the DML7-FAM binding integrin α 3 β 1 receptor protein subunit domain in living cell systems. (A–E) Confocal imaging analysis of DU145 cells including native cells (A), incubation with the saturated DML7-FAM aptamer (B), and subsequent co-incubation with excess saturated G_5 @BSA as a negative control (C), G_5 @anti α 3 (P1B5) (D), and G_5 @anti β 1 (K-20) (E), respectively. (F) FLI analysis of grey values that were estimated using Image J software. Group 1, 2, 3, and 4 indicated the cells in (B), (C), (D), and (E). (G and H) FCM assay on FLI quenching of DML7-FAM induced by different amounts of G_5 @anti α 3 (P1B5) and G_5 @anti β 1 (K-20), respectively. FCM charts included native DU145 cells (cell), incubation with saturated DML7-FAM (+DML7-FAM), and the subsequent co-incubation with G_5 @antibody of different microliters as indicated by the numbers in each panel (++ number). (I) The Φ_{FLI} curves plotted as a function of the amounts of G_5 @anti α 3 (P1B5) (blue data) and G_5 @anti β 1 (K-20) (red data), respectively. (J and K) Linear relationship of F_0/F and $[Q]$ before the saturation binding of G_5 @anti α 3 (P1B5) (J) and G_5 @anti β 1 (K-20) (K), respectively. (L) The calculated R' values between the binding sites of DML7-FAM and each antibody according to the antibody size parameters obtained from TEM (group 1), AFM (group 2) and the theoretical calculation (group 3), respectively.



nonspecific effects of $G_5@antibodies$ (Fig. 8F). Next, FCM was performed to quantitatively determine the initial FLI (I_0) of the saturated DML7-FAM aptamer, quenched FLI(t) in the presence of saturated $G_5@anti\alpha3$ (P1B5) or $G_5@anti\beta1$ (K-20), and the background FLI (blank) of native DU145 cells. The population of the DU145 cells incubated with different ligands all had good monodispersities (Fig. S20A–D†). Next, the quenching effects of DML7-FAM by different amounts of $G_5@anti\alpha3$ (P1B5) or $G_5@anti\beta1$ (K-20) are exhibited in Fig. 8G and H, respectively. A plateau was reached after adding over 200 μL of $G_5@anti\alpha3$ (P1B5) (blue line) or $G_5@anti\beta1$ (K-20) (red line) (Fig. 8I). Accordingly, the stable maximum of Φ values was produced to be nearly 44% by $G_5@anti\alpha3$ (P1B5) and 34% by $G_5@anti\beta1$ (K-20).

According to eqn (4), F_0/F follows a $[Q]$ -dependent linear relationship for both $G_5@anti\alpha3$ (P1B5) and $G_5@anti\beta1$ (K-20), rather than a rising curve bent to the y -axis (Fig. 8J and K), which indicated that the quenching from FLI is dynamic quenching. According to the parameters obtained from TEM, AFM, and theoretical calculations (Table 1), the calculated $R'_{\alpha3}$ value was 14.09 ± 1.54 nm, 14.12 ± 1.45 nm, and 14.10 ± 1.49 nm, respectively, at a stable maximum of Φ in the presence of saturated $G_5@anti\alpha3$ (P1B5) (blue bars, Fig. 8L). Similarly, the calculated $R'_{\beta1}$ value was 16.13 ± 0.72 nm, 16.00 ± 0.69 nm, and 16.08 ± 0.69 nm, respectively, at a stable maximum of Φ in the presence of saturated $G_5@anti\beta1$ (K-20) (red bars, Fig. 8L). Statistical analysis exhibited no significant difference among the three methods, *i.e.*, TEM, AFM, and theoretical calculations, when calculating R' (Fig. S21A and B†). Hence, the data obtained from the three methods were averaged, and the averaged values of $R'_{\alpha3}$ and $R'_{\beta1}$ were 14.1 nm and 16.1 nm, in which the difference was greater than the resolution of the SN-nanoruler and thus had high reliability, *i.e.*, DML7-FAM is much closer to $G_5@anti\alpha3$ (P1B5) rather than $G_5@anti\beta1$ (K-20).

The measurement results of the SN-nanoruler strongly implied integrin $\alpha3\beta1$ acting as a target of the DML7 aptamer and further hints that the binding site of the DML7 aptamer is closer to the $\alpha3$ subunit domain, rather than to the $\beta1$ subunit domain. However, the SN-nanoruler may not accurately pinpoint the binding site of the aptamer, because the working model is highly related to selection of the antibody, and the selection is usually random experimentally. With this in mind, another antibody of the integrin $\alpha3$ subunit, $anti\alpha3$ (VM-2), was examined for competitive binding with DML7-FAM. Coincidentally, a significant competition was observed between DML7-FAM and native $anti\alpha3$ (VM-2) (Fig. S22A and B†). This result not only provided full proof that the integrin $\alpha3$ subunit domain precisely acted as the binding target of DML7-FAM, but also evidenced that the binding site of the DML7 aptamer overlapped with that of $anti\alpha3$ (VM-2) on the integrin $\alpha3$ subunit domain.

Discussion

Our study developed a SN-nanoruler with a single-nucleobase resolution by virtue of a dye-labeled aptamer and single-sized GNP-antibody complex as probes to deduce the molecular distance between two binding ligands at the CD71 protein

subunit domain level on living cell membranes. By lengthening the “ n TA” spacer, *i.e.*, one TA base at a time, between the end of the aptamer and dye, the single-nucleobase resolution of the SN-nanoruler was verified. The SN-nanoruler possessed a high precision, thus exhibiting high potential for *in situ* identification of aptamer-binding sites in living systems, and further advanced the molecular distance measurement down to the receptor protein subunit domain level in living cells. The SN-nanoruler can be a powerful complement or/and replacement to traditional energy transfer nanorulers for dynamically changing molecular event measurement at the high spatial resolution scale in biochemical fields and takes a step forward toward much wider applications of optical-based rulers in observing molecular structure transformation in living systems. For example, the SN-nanoruler is expected to *in situ* project each base of an aptamer on a specific plane, *e.g.*, the GNP surface, in living systems, from the perspective of molecular distance, that enjoy great significance for establishing sensitive, reliable, and convenient aptamer-based diagnostic methods.

Also, our study inspires several key issues that could be explored in the future. (1) The choice of fluorescent dyes is still very limited because of the absence of r_0 values for different dyes and GNP pairs. There is an urgent need to develop novel strategies to easily determine more dyes and GNP pairs' r_0 values, which would carry significance in biochemical sensing. The single-nucleobase resolution-based SN-nanoruler will pave a new way to determine the unknown r_0 value of different GNP-dye pairs in living cell systems. (2) The R' value is defined as the distance between the FAM fluorophore and antibody binding sites, and the fluorophore is labeled on the 5' end of the aptamer. On the molecular scale, the spatial location of the fluorophore should not be identical to that of the aptamer site with specific three-dimensional scaffolds. However, a dye-labelled aptamer is considered a point dipole, which enables the fluorophore location to be the same as the aptamer site experimentally in this paper. Notably, 3'-end modifications of fluorophores in aptamers might produce different results, which could be deduced using the single-nucleobase resolution-based SN-nanoruler as well. (3) Application of the NSET effect is focused on monitoring the ensemble state of the aptamer these days. However, imaging of the transient state, *e.g.*, single events on the living cell membrane, will expand our understanding of how a cell finely regulates molecular events on its membrane.^{42,43} Observing dynamically changing molecular events related to bio/chem-molecules and the transitions within the living biological system is significant to further understanding life processes and diagnosis of disease. In the future, the SN-nanoruler would be a nice spectral measurement tool in the chemo-metrics field, which could be applied for *in situ* monitoring of single event dynamic transitions, bimolecular structure changes, aptamer-protein or protein-protein interaction, cell-cell communication and the “on/off” state of an ion channel.

Data availability

Details of methods, experimental procedures, and characterization data are available in the ESI.† Further information and



requests for resources should be directed to and will be fulfilled by the lead contact, Honglin Liu (liuhonglin@mail.ustc.edu.cn).

Author contributions

Yu Zhang: investigation, conceptualization, methodology, formal analysis, data curation, visualization, writing – original draft, review, and editing. Mengke Su: formal analysis, data curation, writing – review and editing. Xingru Fang: investigation. Wenwen Huang: investigation. Hao Jiang: investigation. Qi Li: investigation. Nisar Hussain: writing – review & editing. Mao Ye: writing – review, editing and supervision. Honglin Liu: conceptualization, methodology, formal analysis, data curation, resources, writing – review, editing and supervision. Weihong Tan: writing – review, editing and supervision.

Conflicts of interest

The authors declare no competing interests.

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