Quantum dot-loaded monofunctionalized DNA Icosahedra for single particle tracking of endocytic pathways

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Abstract

Functionalization of quantum dots (QDs) with a single biomolecular tag using traditional approaches in bulk solution has met with limited success. DNA polyhedra consist of an internal void bounded by a well-defined three-dimensional structured surface. The void can house cargo and the surface can be functionalized with stoichiometric and spatial precision. Here, we show that monofunctionalized QDs can be achieved by encapsulating QDs inside DNA icosahedra and functionalizing the DNA shell with an endocytic ligand. We deployed the DNA-encapsulated QDs for real time imaging of three different endocytic ligands - folic acid, galectin-3 (Gal3) and the Shiga toxin B-subunit (STxB). Single particle tracking of Gal3 or STxB-functionalized, QD-
loaded DNA icosahedra allows us to monitor compartmental dynamics along endocytic pathways. These DNA-encapsulated QDs that bear a unique stoichiometry of endocytic ligands represent a new class of molecular probes for quantitative imaging of endocytic receptor dynamics.

The ubiquitous deployment of quantum dots (QDs) in biology has been constrained due to lack of methodologies to permanently and homogenously monofunctionalize them. Traditional approaches to achieve this involve coupling QDs and ligands in fixed ratios in solution (Supplementary table 1). This has met with limited success due to stochastic control over ligand stoichiometry leading to sample heterogeneity.

Structural DNA nanotechnology has yielded diverse, well-defined, nanoscale polyhedra. These polyhedra enclose an internal void that can house nanoscale cargo and possess a well-defined surface for molecular display with both stoichiometric and spatial precision. DNA icosahedra can encapsulate cargo without compromising cargo functionality and can be targeted to specific endocytic pathways. Here, we combine the photostability of QDs, with the molecular programmability of the DNA polyhedra by encapsulating QDs inside DNA icosahedra displaying a solitary bio-molecular tag. This is a new class of precisely functionalized particles of homogenous stoichiometry for long duration live imaging.

We first pinpoint specific residues on the icosahedron for optimal surface display of biological tags. We then encapsulate QDs in the icosahedron and monofunctionalize the scaffold with folic acid (FA), galectin-3 (Gal3), or Shiga toxin B-subunit (STxB). The stability of QDs by previous approaches is limited in that ligands on the QD surface are labile and can leach off leading to loss of the tag or QD aggregation. In our approach, this problem is overcome since the issue of QD surface chemistry is completely transferred to the robust, controllable chemistry of DNA. Monofunctionalization is achieved by conjugating the bio-tag to an amine group displayed on the DNA shell without engaging the QD. This effectively results in stably monofunctionalized QDs. Using DNA-encapsulated QDs monofunctionalized with Gal3 or STxB as endocytic ligands, we track plasma membrane binding, bending, intracellular uptake and long duration dynamics of endocytic carriers on the Gal3/STxB pathways.

**Encapsulation of QDs within DNA icosahedra**

CdSe/CdS/ZnS-based quantum dots of 5 nm diameter (QD5) were synthesized as described (Supplementary Fig. S1). DNA icosahedra, I, were assembled from three types of five-way junctions (5WJs), V, U and L8. To encapsulate QDs within DNA icosahedra, two half-icosahedra VU5 and VL5 were incubated in the presence of excess of QD5 (Fig. 1a and methods). QD-loaded DNA icosahedra were purified by gel electrophoresis and size-exclusion chromatography, (SEC-HPLC; Fig. 1b-c). A size fraction corresponding to the DNA icosahedra showed fluorescence corresponding to QD5, indicating the formation of an I-QD5 complex (I\textsubscript{QD}) (Supplementary Fig. S2,6). Dynamic light scattering (DLS) studies revealed that free QD5 showed an R\textsubscript{H} of 4.5±1.8 nm, while I showed an R\textsubscript{H} of 9.5±0.1 nm (Fig. 1d, Supplementary Fig. S3). Purified I\textsubscript{QD} showed an R\textsubscript{H} of 10.6±0.4 nm (Fig. 1d). This implies that QD5 in I\textsubscript{QD} associates with the DNA icosahedron such that it does not
significantly alter the icosahedron dimensions. For an electron dense particle such as QD5 this is possible only if QD5 is encapsulated within the DNA icosahedron.

Since the DNA icosahedron has a pore size of ~2.5 nm, we used a differential quenching assay to test whether the QDs were encapsulated. Free QD5 and I_{QD} were subjected to quenchers of sizes ranging from 0.5-5 nm. The fluorescence intensities of 5 nM free QD5 and I_{QD} were measured in the presence of fixed amounts of each quencher. When treated with 1/K_{SV} concentrations of each quencher, quenchers of all sizes quenched QD5 fluorescence intensity by 50% (Fig. 1e, Supplementary Fig. S4). However, for I_{QD}, only quenchers below ~2.2 nm diameter quenched its fluorescence by 50%. Quenchers ≥ 3 nm could not quench I_{QD} fluorescence at all, while quenchers between 2.2-3 nm partially quenched the fluorescence (Fig. 1e). This confirms that QD5 in I_{QD} is physically encapsulated as cargo within the DNA icosahedron. This was reaffirmed by experiments with single QDs or single I_{QD} performed on a confocal microscope (Fig. 1f, Supplementary Fig. S5 & S6).

QDs with sizes from 5 to 11 nm, with different surface chemistries, could be encapsulated in DNA icosahedra indicating the generalizability of this method. The capacity of DNA icosahedra to encapsulate nanoparticles was demonstrated by transmission electron microscopy (TEM) where QDs were substituted with 10 nm gold nanoparticles (GNP10) (Fig. 1g, Supplementary Fig. S7). The sizes of the DNA shell were between 15-25 nm, consistent with previous TEM measurements of DNA icosahedra.

**Endocytic uptake maps nucleotide positions on the DNA Icosahedron**

To target QD-loaded DNA cages along specific endocytic pathways, we first identified those nucleotide positions on the DNA scaffold where a conjugated molecular tag faces the external milieu. We developed an atomistic model of the DNA icosahedron conjugated to folic acid (FA) using xLeAP module in AMBER and studied its *in silico* stability using molecular dynamics (MD) simulations (see methods). After 50 ns of simulation, the structure displayed moderate deviations along the edges and minor structural changes at the 5WJs. However, it retained its icosahedral geometry indicating the overall stability of the architecture *in silico* (Fig. 2a-b).

We then conjugated a small endocytic ligand, folic acid (FA) to different nucleotide positions along a given edge of the DNA icosahedron and quantitated FA accessibility by endocytic uptake in cells expressing the folate receptor. Seven different folate-conjugated icosahedra (I_{FA}) were realized, each with a single folate tag at residues 7, 9, 11, 13, 15, 17, and 19 away from the vertex V, collectively spanning a full helical turn (Supplementary Fig. S8a,b). An Alexa 647 fluorophore was also incorporated on I_{FA} to give I_{FA/A647} in order to visualize the icosahedral shell. IA2.2 cells stably expressing the human transferrin and folate receptors, were pulsed with 100 nM of each of the seven distinct I_{FA/A647} samples along with 100 nM fluorescent transferrin (TfA568) to normalize for endocytic uptake. I_{FA/A647} samples with FA tags located at positions 11 and 13 showed maximal endocytic uptake that decreased sharply as the position of the tag was moved along either direction (Fig. 2c). This is consistent with the atomistic model where, at positions 11 and 13 the FA tag is maximally
exposed, reaffirming its enhanced accessibility to the folate receptor (Supplementary Fig. S18).

We then tested whether \( I^{\text{FA/A647}} \) was endocytosed specifically by the folate receptor pathway (Fig. 2d-e). \( I^{\text{FA/A647}} \) with FA at position 11, hereafter designated as \( I^{\text{FA/A647}} \), showed efficient uptake and colocalization with endocytic markers Tf (Fig. 2d, upper panels) and pteroyl lysine bodipy-TMR (PLB), a fluorescent analog of folic acid (Fig. 2d, lower panels). When \( I^{\text{FA/A647}} \) was co-pulsed with 10 fold excess free FA, it was successfully competed out (Fig. 2f, upper panels). \( I^{\text{A647}} \) without an FA tag also showed no uptake. When IA2.2 cells lacking the folate receptor were pulsed with \( I^{\text{FA/A647}} \) no uptake was observed (Fig. 2f, lower panels). This indicates that uptake of \( I^{\text{FA/A647}} \) needs the folate tag and occurs via the folate receptor pathway (Fig. 2f-g). Further, the internalization efficiency showed a sinusoidal pattern as a function of nucleotide position, with a periodicity that matches the pitch of a B-DNA helix (Figure 2c). This proves monofunctionalization of DNA architectures in bulk solution.

Given that monofunctionalized DNA icosahedra mark a specific endocytic route, we combined this with \( I^{\text{QD}} \) bearing an FA tag at position 11 (\( I^{\text{QD/FA}} \)). \( I^{\text{QD/FA}} \) was efficiently uptaken and colocalized with Tf\( ^{\text{A488}} \) showing specific targeting of \( I^{\text{QD/FA}} \) along the folate pathway (Fig. 2h,i). Thus we could realise molecularly identical \( I^{\text{QD}} \)s displaying a single ligand on the icosahedral DNA shell accessible to its cognate endocytic receptor.

### Binding and endocytic uptake of \( I^{\text{QD/Gal3}} \) and \( I^{\text{GNP/Gal3}} \)

We confirmed monofunctionalization as well as expanded the range of conjugated endocytic tags by probing the stoichiometry of a cellular lectin, galectin-3 (Gal3), conjugated to the DNA icosahedron (Fig. 3a, Supplementary Fig. 9). DNA icosahedra bearing a surface displayed amine group (\( I^{\text{NH2}} \)) were each loaded with a solitary 10 nm gold nanoparticle (GNP10) to give \( I^{\text{GNP/NH2}} \), as previously described. His-tagged Gal3 bearing an engineered cysteine residue (His-Gal3/Cys) was conjugated to \( I^{\text{GNP/NH2}} \) and purified to give \( I^{\text{GNP/His-Gal3}} \) (Supplementary Fig. S9a,b). To \( I^{\text{GNP/His-Gal3}} \) deposited on a TEM grid, 5 nm gold nanoparticles bearing Ni-nitrilotriacetate (GNP5) was added. When visualized by TEM, 74% of 10 nm GNPs were present in close proximity with a GNP5 (Fig. 3b-d). We also observed 23% of solitary 10 nm GNPs, probably due to \( I^{\text{GNP/His-Gal3}} \) orientations on the grid where the Gal3 tag is inaccessible to GNP5. 3% of \( I^{\text{GNP/His-Gal3}} \) showed more than one proximal GNP5. Indeed, pure GNP5 solutions show ~30% dimers/oligomers, which cannot be abolished even with strong sonication and dilution (Supplementary Fig. S10). Non-functionalized \( I^{\text{GNP}} \) shows no such paired association with GNP5 (Supplementary Fig S10). Thus, single particle TEM analysis revealed that the robust generation of monofunctionalized, cargo-loaded DNA icosahedra in high yields (65-90%; Fig. 3c).

Given the exquisite control over mono-functionalization, we tested whether \( I^{\text{QD/Gal3}} \) could be applied to transport QDs or GNPs intracellularly and probe the Gal3 endocytic pathway. Gal3 binds glycosylated proteins such as \( \beta1 \) integrin and CD44 that are resident on the plasma membrane of cells. Gal3 then undergoes oligomerization driving biogenesis of morphologically distinct crescent-shaped clathrin-independent carriers (CLICs) mediated by
glycosphingolipids. CLICs eventually fuse with early/sorting endosomes. Using sulfo-
MBS chemistry, Gal3/Cys was conjugated to IQD to give IQD\textsuperscript{Gal3}. We tested the cellular
binding and uptake pathway of IQD\textsuperscript{Gal3} by colocalization with fluorescently labeled Gal3
(Gal3\textsuperscript{Cy5})(Fig. 3e, Supplementary Fig. S9a-c). IQD\textsuperscript{Gal3} and Gal3\textsuperscript{Cy5} bound the plasma
membrane of mouse embryonic fibroblasts (MEFs) efficiently with quantitative
colocalization (Fig. 3e). In the presence of a specific Gal3 competitor such as lactose (100
mM), binding of both IQD\textsuperscript{Gal3} and Gal3\textsuperscript{Cy5} was reduced by 95% indicating that binding was
specific (Fig. 3f). Upon depleting cellular ATP, the scission of Gal3-induced membrane
invaginations is inhibited leading to distinctive, long, tubular plasma membrane
invaginations. When IQD\textsuperscript{Gal3} and Gal3\textsuperscript{Cy5} were incubated with ATP-depleted cells on ice
for 30 min, both IQD\textsuperscript{Gal3} and Gal3\textsuperscript{Cy5} colocalized in long, tubular plasma membrane
invaginations (Fig. 3g). Thus IQD\textsuperscript{Gal3} retains its binding specificity to the cell membrane and
induces downstream plasma membrane invagination.

Gal3-induced CLICs are distinctive, crescent-shaped, short, tubular structures as revealed
by TEM. To unequivocally prove that IQD\textsuperscript{Gal3} is internalized by the Gal3 pathway, we
sought to visualize Gal3 within such CLICs. To facilitate visualization by TEM, we created
GNP\textsuperscript{Gal3}, encapsulating 5-6 GNPs of ~5 nm diameter. MEFs were incubated with Gal3
conjugated to horseradish peroxidase (Gal3-HRP) and GNP\textsuperscript{Gal3} in a 1:1 ratio and HRP was
developed using 3,3'-diaminobenzidine-H\textsubscript{2}O\textsubscript{2}. TEM (see methods) images of these cells
clearly revealed crescent shaped structures near the plasma membrane - the definitive
signature of CLICs (Fig. 4a-c). Under low contrast conditions, CLICs containing clusters of
5 nm GNPs could clearly be seen, indicating that GNP\textsuperscript{Gal3} and Gal3-HRP are endocytosed
together into CLICs (Fig. 4b-c, right panels). From two independent experiments,
comprising 25 cells, 94% of GNPs localized in DAB precipitate-positive structures (Fig.
4b,c), confirming that GNP\textsuperscript{Gal3} adopts the same pathway as Gal3. Consistent with literature,
78% of the intracellular structures had CLIC morphology.

To address the endocytic fate of these particles, IQD\textsuperscript{Gal3} and Gal3-Cy5 were co-incubated
with MEFs for 2, 15 and 60 min (Fig. 4d, Supplementary fig. S15). At 2 min, IQD\textsuperscript{Gal3} and
Gal3-Cy5 colocalized in small vesicular structures close to the plasma membrane (Fig. 4d,
upper panel). At 15 min, they colocalized in larger punctate structures, likely corresponding
to early/sorting endosomes (Fig. 4d, middle panel). At 60 min, IQD\textsuperscript{Gal3} and Gal3-Cy5
colocalized in large, ring-like, perinuclear structures, likely corresponding to late endosomes
(Fig. 4d, lower panel). This quantitative colocalization with Gal3-Cy5 along the entire
pathway (Pearson’s coefficient > 0.8, Spearman’s coefficient > 0.9 and Mander’s coefficient
> 0.9 at all time points), confirmed that post attachment of Gal3, IQD does not alter Gal3
internalization characteristics.

**Single molecule tracking of IQD\textsuperscript{STxB} uptake in live cells**

The B-subunit of Shiga toxin (STxB) is a homopentameric protein that binds the
glycosphingolipid Gb3 on the plasma membrane of particular eukaryotic cells. Post binding,
STxB pentamers cluster to drive the formation of membrane invaginations that undergo
dynamin, endophilin-A2 and actin-dependent scission to form endocytic carriers that then
fuse with early endosomes, followed by further trafficking along the retrograde
route36. An STxB variant carrying an engineered cysteine residue at its C-terminal end was coupled to monofunctionalized IQDs, as described earlier (Supplementary figs S11-13) to give IQD^{STxB}. Alexa488 labeled STxB (STxB^{A488}) and IQD^{STxB} efficiently bound the plasma membrane, were internalized into punctate structures and colocalized with each other (Fig. 5a,b). Further, IQD^{STxB} localized in early endosomes (EE) via the STxB internalization pathway (Supplementary Fig. S14 & S17).

Given the specific uptake of IQD^{STxB} and photostability of QDs, we investigated the dynamics of STxB-mediated endocytic transport by real-time TIRF microscopy, with single particle precision (Supplementary Fig. S16). HeLa cells were pulsed with a mixture of ~200 nM unlabelled STxB doped with 50 pM IQD^{STxB}. This induced efficient clustering of STxB on the membrane that incorporated IQD^{STxB} within larger STxB clusters. Upon internalisation, IQD^{STxB} disappeared from the plasma membrane plane of observation (Fig. 5c, left panel, Supplementary movie S1). Before internalization, most IQD^{STxB} particles undergo hop diffusion characterized by motion in confined spaces followed by sudden jumps (Fig. 5c, middle and right panels, Supplementary movie S2). IQD^{STxB} diffusion on the extracellular leaflet of the plasma membrane therefore follows a complex diffusion behaviour that suggests picket-fence type compartmentalization by the underlying actin meshwork, as seen in phospholipids37. Compartment sizes characterized by the confined diffusion ranged from 30-80 nm with average diffusion coefficients of 0.12 µm²/s, similar to those observed for transferrin receptors38.

Endosomes containing IQD^{STxB} colocalized with and moved along microtubules actively, (Fig. 5d, Supplementary movie S3), transferring back and forth between microtubules (Supplementary movie S4). Endosomes showed bursts of active transport with pause intervals indicating either normal or confined diffusive behaviour respectively (Fig. 5e). A single endosome switches between these two modes, with active bursts showing run lengths of 2.4 ± 1.4 µm and average speeds of 0.5 ± 0.2 µm/s. We used mean square displacement analysis (MSD) from where α values were used to characterize the spectrum of motion observed for a population of endosomes. A measure of the scaling exponent, α < 0.4 indicates confined motion, 0.4 < α < 1.45 indicates diffusive motion, while α > 1.45 indicates directed motion. The observed distribution of α values in our data (see SI, methods) reveals that a majority of endosomes, at any given time display either confined or diffusive behaviour (Fig. 5e). This average value is similar to velocities observed in previous studies of Rab5 positive endosomes, suggesting that most of the motile IQD^{STxB} observed are in early endosomes39. Stochastic active bursts result in the overall transport of endosomes. The active motions also showed reversals, suggesting that both plus and minus end microtubule-associated motors are present on the endosomes. The observed pauses may arise from several phenomena: endosomes crossing over at intersections of microtubules, interaction with ER, actin meshwork or other organelles40,41. Endosomes containing IQD^{STxB} could be categorized into static early endosomes (SEE) and dynamic endosomes (DE), as observed previously for transmembrane cargo and viral particles42. Internalized IQD^{STxB} actively moved and fused with SEE (Fig. 5f, yellow arrows). Small vesicles containing IQD^{STxB} emerge from these SEE and then move actively, constituting dynamic endosomes (DE) (Fig. 5f, white arrows). The broad range of α values reflects the crowded nature of the intracellular milieu and the interaction of IQD^{STxB} containing endosomes with
various intracellular structures such as actin meshwork or the microtubule network in between directed active runs.39.

Conclusions

We show the efficient encapsulation of quantum dots (QDs) within a DNA icosahedron in bulk solution. Encapsulation does not alter QD fluorescence properties in vitro or in the cellular milieu. Endocytic uptake assays supported by molecular modeling studies pinpointed residues on the icosahedron that display biological tags most efficiently. By site-specifically monofunctionalizing these icosahedra with endocytic tags we could realize QDs homogenously bearing a single targeting ligand (folate/Gal3/STxB). This enabled the live tracking of long duration compartment dynamics in cells. This methodology is generalizable across both QDs and endocytic tags, due to the capsular DNA interface. As the icosahedral DNA surface is well defined, one can envisage homogenous oligofunctionalization of IQDS with ligands. These could probe receptor-ligand interactions and oligomerization, whose nature and consequences change with ligand-receptor stoichiometries.

Methods

Materials

The unlabeled and modified, labeled oligonucleotides (HPLC-purified and lyophilized) were obtained from IBA GmbH. N-cyano imidazole (NCI) was synthesized in-house according to previous protocols. l-octadecene (ODE, Sigma), oleylamine (Sigma), oleic acid (Sigma), trioctylphosphine (TOP, Cytec), cadmium oxide (Sigma), selenium pellets (Sigma), sulfur powder (Sigma) and tetradecylphosphonic acid (TDPA, PCI synthesis) were purchased from the indicated suppliers. Methods for DNA icosahedron construction and characterization have been previously described. CdSe/CdS/ZnS QDs were synthesized according to previously published protocols. The detailed procedures for synthesis, characterization and encapsulation of QDs within DNA icosahedron are provided in the supplementary information. The experiments on cells were carried out using QDs provided by Nexdot (www.nexdot.fr).

Cell Culture and labeling with endocytic markers

IA2.2 cell line is a Chinese hamster ovary (CHO) cell line that lacks endogenous transferrin receptors but stably expresses the human transferrin and folate receptors described in reference 43. The cells were cultured in Ham’s-F12 Complete media (HF-12, Himedia) containing 10% heat-inactivated FBS, 100 μg/mL streptomycin and 100 μg/mL penicillin with 200 μg/mL G418 and 100 μg/mL hygromycin to ensure maintenance of the transferrin and folate receptors. HeLa (source described in reference 44), Rab5-GFP expressing HeLa cells and MEFs cells (source described in ref 31) were cultured in DMEM media supplemented with 10% FCS and PS mixture. For binding experiments, IA2.2 cells were pulsed for 30 min on ice with 100 nM concentrations of all probes (IQDFA, Tf or PLB), washed 2-3 times with M1 media, and imaged under Olympus FV1000 confocal microscope, using appropriate lasers and imaging conditions. For the uptake assay, cells were pulsed for 15 min at 37°C with the indicated probes, washed and imaged. Uptake and
colocalization were quantified using the Pearson colocalization coefficient application in ImageJ (NIH). His-tagged cysteine engineered Gal3 was purified according to previously established protocol34. Cysteine engineered Gal3 and STxB were conjugated to amine modified DNA using sulfo-MBS or SM(PEG)8 (PierceNet) crosslinkers. Labeled Gal3/STxB and DNA-Gal3/DNA-STxB were pulsed to cells at 50 or 100 nM concentrations in DMEM media for the indicated periods of time. Post incubation cells were washed with PBS, fixed in 4% PFA, and coverslips mounted on fluoromount Mowiol for imaging. For fixed samples, imaging was performed using a Leica epifluorescence microscope, and for live cell and single particle tracking, A1R confocal, TIRF-FRAP and spinning disk SP7 microscopes from Nikon were used. The detailed procedures are provided in supplementary information.

**Supplementary Material**

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Fig. 1. Encapsulation of quantum dots (QDs) within DNA icosahedra (I).

a, (left) Schematic showing the formation of quantum dot-loaded icosahedra (I_{QD}). Two complementary half icosahedra VU_5/VL_5 are mixed in a 1:1 ratio in the presence of excess QDs, purified from free QDs to get I_{QD}. (right) Cartoon representation of all molecular tags used to functionalize the DNA icosahedron.

b, Gel electrophoretic mobility shift assay for the formation of I_{QD}. Fluorescence image of 0.8% agarose gel in 1X TAE (λ_ex at 488 nm): Lane 1, QD (λ_em = 605 nm); lane 2, VU_5FITC (λ_em = 520 nm), lane 3, I_{QD}FITC (λ_em = 520 & 605 nm), lane 4, I_{FITC} (λ_em = 520 nm).

c, Size-exclusion chromatogram of I_{QD} post-gel excision where absorbance at 260 nm was followed.

d, Dynamic light scattering traces of free QD (green), I (black) and I_{QD} (red).

e, Fluorescence intensity-based quenching assay for free QD (green) and I_{QD} (red) in bulk solution. Quenchers are gold nanoparticles (GNPs) of indicated sizes, iodide (0.5 nm), TEMPO (1 nm) and TEMPO Dextran (2.5 nm). Mean values of two experiments are presented with corresponding s.d. (n=2).

f, Single molecule quenching assay for free QD and I_{QD} when subjected to 2 nm and 5 nm size GNPs. Error bars indicate mean of two experiments with associated s.d. (two-tailed unpaired
t-test. * $p < 0.0001$, $n=2$.

Representative TEM image of GNP-encapsulated DNA icosahedra stained with 1% uranyl acetate. Scale bar: 200 nm.
Fig. 2. Cellular validation of atomistic model of DNA icosahedron.

a-b, Atomistic model of charge neutralized, solvent stabilized DNA icosahedron post 50 ns of MD simulation showing a, the C3 and b, the C5 axes of symmetry. c, Cellular uptake of IFA/A647 as a function of different nucleotide positions of FA on IFA/A647. Uptake is normalized with respect to TfA568 as an internal control. Mean values of total cell intensity from three independent experiments. d-g, Uptake of IFA/A647 is through the folate receptor pathway. d, Colocalization of endosomes containing IFA/A647 (red) uptaken by IA2.2 cells with endocytic probes (green) transferrin (TfA568) and pteroyllysine (PLB). e, Quantification
of colocalization of $I^{FA/A647}$ with $Tf^{A568}$ and PLB. Mean Pearson’s correlation coefficient (PCC) for $n = 20$ cells with associated s.d. ($n=2$) and for images shifted by 10 pixels. $f$, $I^{FA/A647}$ uptake occurs through the FA tag and the folate receptor. $I^{FA/A647}$ uptake in the presence of 10-fold excess FA (upper panels). $I^{FA/A647}$ uptake in TRVb-2 cells lacking the folate receptor (bottom panels). $g$, Quantification of $I^{FA/A647}$ uptake in $f$. Mean values of two independent experiments with associated s.d. $h,i$, QDs encapsulated in $I^{FA}$ ($I_{QD}^{FA}$) shows the same uptake pathway as $I^{FA/A647}$. $h$, Cellular uptake of Tf$^{A647}$ (red) and $I_{QD}^{FA}$ (green) in IA2.2 cells post 15 min incubation at 37°C. $i$, Endocytic uptake of Tf$^{A647}$ and $I_{QD}^{FA}$ quantified for 15 cells. AF = autofluorescence. All scale bars are 10 µm, inset scale bars are 5 µm. All error bars are standard errors and use the two-tailed unpaired t-test.
Fig. 3. Binding of I$_{QD}^{Gal3}$ to the plasma membrane of cells:

a. Schematic of the assembly strategy to tag I$_{QD}$ with galectin-3 (Gal3) to give I$_{QD}^{Gal3}$. U is conjugated to Gal3 and then incorporated into VU$_5$ to give VU$_5^{Gal3}$. Assembly of VU$_5^{Gal3}$, VL$_5$ and QDs yields I$_{QD}^{Gal3}$. b, TEM images showing that DNA icosahedra each encasing a single 10 nm gold nanoparticle (GNP) is functionalized with a single Gal3 tag. I$_{GNP}^{His-Gal3}$ incubated with excess of 5 nm NTA-coated GNPs (GNP5) imaged by TEM. Scale bar: 50 nm. c, Frequency of the number of GNP5 particles within a 10 nm radius of I$_{GNP}^{His-Gal3}$ particles d, Zoomed images of representative examples of I$_{GNP}^{His-Gal3}$ attached to a single
GNP5. e,f, I_{QD}^{\text{Gal3}} and Gal3 show similar plasma membrane binding characteristics. e, Fluorescence images of mouse embryonic fibroblasts (MEFs) incubated at 4°C with I_{QD}^{\text{Gal3}} (green) and Gal3^{Cy5} (red) f, Binding of both I_{QD}^{\text{Gal3}} and Gal3 is competed out by 100 mM lactose. g, Fluorescence images of I_{QD}^{\text{Gal3}} (green) and Gal3^{Cy5} (red) on ATP-depleted MEFs. Insets show tubular membrane invaginations induced by Gal3 due to ATP-depletion. All scale bars: 10 µm. All inset scale bars: 1 µm
Fig. 4. TEM studies reveal that I$_{QD}^{Gal3}$ is endocytosed through clathrin-independent carriers (CLICs).

a-c, I$_{GNP}^{Gal3}$ is present in CLICs.  

a, TEM sections of MEFs pulsed with Gal3-HRP for 3 min at 37°C and developed with diaminobenzidine/H$_2$O$_2$ (left panel). Zoomed image of a Gal3 containing CLIC, showing crescent shaped morphology (see arrowhead, right panel).

b-c Left panels show representative TEM sections of MEFs pulsed with 1:1 Gal3-HRP : I$_{GNP}^{Gal3}$ at high contrast showing crescent-shaped CLICs (boxed areas) Scale bar: 1 µm. Right panels show zoomed TEM images of boxed regions in the left panels at low contrast.
revealing GNPs in the CLICs (arrowheads). Scale bar: 100 nm. d-f. I\textsubscript{QD}\textsuperscript{Gal3} follows the Gal3 endocytic route. d, Colocalization of I\textsubscript{QD}\textsuperscript{Gal3} (green) and Gal3\textsuperscript{Cy5} (red) at the indicated chase times in MEFs: d, 2 min in CLICs, e, 15 min in early endosomes f, 60 min in late endosomes. Scale bar: 10 µm; inset scale bar: 1 µm.
Fig. 5. Single particle tracking of IQD\textsuperscript{STxB} in live cells.

a-b, IQD\textsuperscript{STxB} adopts the STxB endocytic route. a, STxB\textsuperscript{A488} (green) and IQD\textsuperscript{STxB} (red) bind the plasma membrane of HeLa cells. b, STxB\textsuperscript{A488} (green) and IQD\textsuperscript{STxB} (red) colocalize at 2 min chase time. Inset shows zoomed image of the labelled compartments.

c, Single particle tracking of IQD\textsuperscript{STxB} bound on the plasma membrane of HeLa cells observed using total internal reflection fluorescence (TIRF) microscopy (see Supplementary movie S1). (i) TIRF image of individual IQD\textsuperscript{STxB} particles bound on the section of the plasma membrane. (ii) A collection of single particle tracks of individual IQD\textsuperscript{STxB} particles, differently colored for clarity, obtained from the region shown in (i) (Supplementary movie S2). (iii-iv) Two typical single particle tracks showing (iii) confined diffusion and (iv) a directed run, both color coded for increasing velocities in µm/s.

d-e, Single particle tracking of early endosomes containing IQD\textsuperscript{STxB} moving along microtubules in HeLa cells. d(i) Spinning disc confocal image of early endosomes containing IQD\textsuperscript{STxB} (white) in HeLa cells expressing tubulin-GFP.
(Supplementary movie S3). (ii) Confocal image of early endosomes labeled with I\DOTSTxB (red) localized on microtubules (green), indicated by arrowheads in the merged image. e, A plot of frequency of alpha values (α) obtained from ~ 5470 tracks (grey trace). Also shown are the frequency of trajectories showing confined/diffusive behavior (blue trace) and active directed runs (red). The graph is divided into three sections: light blue, yellow and light pink for confined behaviour, diffusive behaviour, and directed runs respectively. Shown on top are typical tracks for each type of behaviour. f, Montage of a typical I\DOTSTxB labeled static early endosome or sorting endosome (SEE). The white arrowhead indicates a fission event that gives rise to a smaller dynamic endosome (DE). The yellow arrowhead indicates a fusion event of a vesicle arising from the plasma membrane (PM) with the SEE. All scale bars are: 10 µm. Insets, c(iii), c(iv), d(i) and f scale bars: 1 µm.