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Virus-Inspired Membrane Encapsulation of DNA Nanostructures To Achieve In Vivo Stability

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ABSTRACT DNA nanotechnology enables engineering of molecular-scale devices with exquisite control over geometry and site-specific functionalization. This capability promises compelling advantages in advancing nanomedicine; nevertheless, instability in biological environments and innate immune activation remain as obstacles for in vivo application. Natural particle systems (i.e., viruses) have evolved mechanisms to maintain structural integrity and avoid immune recognition during infection, including encapsulation of their genome and protein capsid shell in a lipid envelope. Here we introduce virus-inspired enveloped DNA nanostructures as a design strategy for biomedical applications. Achieving a high yield of tightly wrapped unilamellar nanostructures, mimicking the morphology of enveloped virus particles, required precise control over the density of attached lipid conjugates and was achieved at 1 per ~180 nm². Envelopment of DNA nanostructures in PEGylated lipid bilayers conferred protection against nuclease digestion. Immune activation was decreased 2 orders of magnitude below controls, and pharmacokinetic bioavailability improved by a factor of 17. By establishing a design strategy suitable for biomedical applications, we have provided a platform for the engineering of sophisticated, translation-ready DNA nanodevices.

KEYWORDS: DNA - nanotechnology - lipid bilayer - in vivo - nanostructure - imaging - immune - pharmacokinetics - biodistribution - PEG

Biomedical nanotechnology has undergone considerable progress over the past few decades, from an initial demonstration of liposomes1,2 to that of a logic-gated nanorobot.3 Much of this development has focused on improving the detection and treatment of cancer by increasing the bioavailability and targeting specificity of anticancer agents.4–11 The rapidly growing field of structural DNA nanotechnology12–16 could advance these aims by expanding the range of available nanoparticle geometries and improving the precision of ligand functionalization, two key design parameters. Even more exciting is DNA nanotechnology’s potential for engineering of highly sophisticated nanoscale devices. Recent studies include the nanorobot,3 programmable immuno-adjuvants,17,18 a synthetic membrane channel,19 and a molecular cascade capable of autonomously processing multiple inputs to determine cell phenotype.20 Translation of such devices into biomedical applications requires molecular engineers to first address the susceptibility of DNA nanostructures to nuclease degradation,21 as well as their ability to activate an inflammatory immune response.17,22 Inspired by the stability provided to enveloped viruses via their lipid membranes, we developed enveloped DNA nanostructures that address the above challenges (Figure 1a).

RESULTS AND DISCUSSION

To mimic the geometry of a viral protein capsid shell, we designed a wireframe DNA nano-octahedron (DNO) with an estimated diameter of ~50 nm (Supporting Information Figure 1).23 The octahedron struts are each composed of a bundle of six ~28 nm long double helices14,24 engineered with a ~90° curvature (see Supporting Information Note 1).15 DNOs were self-assembled in a one-pot reaction by combining phage-derived scaffold DNA (p7308)25 with 144...
oligonucleotide staple strands (see Supporting Information Table 1) in a 15 h thermal annealing ramp. Correctly folded structures were isolated by glycerol–gradient centrifugation and verified by negative-stain transmission electron microscopy (TEM) (Figure 1b and Supporting Information Figure 2).

Functional molecular features coupled to oligonucleotides can be assembled onto DNA nanostructures with high precision through hybridization to single-stranded DNA “handles” designed into the nanostructure. We designed an “inner” set of 12 identical handles (protruding from the inside face of the DNO struts) for attachment of fluorophore-conjugated oligonucleotides, providing optical contrast agent functionality. We designed a second “outer” set of 48 handles (protruding from the outside face) for attachment of lipid-conjugated oligonucleotides with the lipid positioned ∼5 nm (2 nm 6-thymidine spacer + 2.5 nm double helix width + 0.5 nm linker for lipid) from the nanostructure frame to drive tight wrapping of the membrane around the structure.

Our encapsulation strategy involves directing lipid bilayer assembly around DNO, recruited by individual lipid-conjugated oligonucleotides preassembled onto outer handles. The lipid bilayers are reconstituted out of a solution of mixed lipid and surfactant.

Figure 1. Schematic of the encapsulation strategy and negative-stain TEM. (a) Lipid–DNA conjugates are annealed to outer handles of non-encapsulated DNA nano-octahedron (N-DNO) in a surfactant solution that forms micelles around the conjugates. Liposomes are added, resulting in mixed surfactant–lipid micelles. Dialysis selectively removes the surfactant and results in a fused lipid bilayer around the DNO. (b) TEM images of purified N-DNO, (c) E-DNO, showing a tightly wrapped unilamellar membrane around the nanostructures, and (d) phi12 bacteriophage, showing comparable ultrastructure and dimensions to E-DNO (scale bar = 50 nm).
(N-octyl-β-D-glucopyranoside) through a dialysis step that selectively removes the surfactant\(^{28}\) to achieve the desired encapsulated DNO (E-DNO) (Figure 1c). We refer to DNO that has not been treated with lipids as non-encapsulated DNO (N-DNO). Unless otherwise stated, our membrane formulation includes 1,2-dioleoyl-sn-glycero-3-phosphocholine (DOPC) (94.2% molar contribution), 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine-N-[methoxy(polyethylene glycol)-2000] (PEG-DOPE) (5%), and fluorescent rhodamine-1,2-dioleoyl-sn-glycero-3-phosphoethanolamine (Rh-DOPE) (0.8%).

We measured internalized DNO diameters of 53 nm versus outer membrane diameters of 76 ± 4 nm from TEM images of E-DNO (Supporting Information Figure 3). Our membrane formulation has an estimated thickness of ∼4 nm lipid bilayer plus ∼5 nm of PEG on each leaflet; therefore, the observed differential of ∼12 nm per vesicle side is consistent with a unilamellar envelope. As anticipated based on our design inspiration, the ultrastructure and dimensions of E-DNO are similar to enveloped virus particles (e.g., phi12 bacteriophage, Figure 1d\(^{29}\)) with our nanostructure taking the place of the protein capsid shell, wrapped tightly by a single lipid bilayer.

To optimize our bilayer reconstitution strategy, we synthesized additional variants of DNO having 0, 12, or 24 outer handles. After subjecting these to the lipid treatment described above, we examined the extent of association between nanostructures and vesicles using a membrane-impermeable DNA dye (PicoGreen). Fluorescence-based quantification of DNO before and after membrane destabilization with surfactant revealed a direct correlation between outer handle number and the percent of nanostructures inaccessible to the dye (Figure 2a). We observed 20.2 ± 3.5% (±SEM) of the 0 handle variant was membrane-enclosed, suggesting there is a nonspecific interaction between the DNA and lipids. This could be tuned by addition of negatively charged lipid (15% 1,2-dioleoyl-sn-glycero-3-phospho-L-serine) to the membrane formulation, which decreased the nonspecific interaction to 5.1 ± 2.7% (Supporting Information Figure 4). The variant with 12 outer handles at an average density of ∼1 handle per 710 nm\(^2\) of nanostructure surface (surface area is defined as that of a sphere enclosing the DNO) showed 43.9 ± 3.1% membrane-enclosed. A comparable 48.3 ± 3.5% was measured with 24 handles (1 handle per ∼350 nm\(^2\)). The association between DNO and membranes was strongest with our design maximum of 48 outer handles (1 handle per 180 nm\(^2\)/handle), which showed 68.1 ± 7.3% enclosed within membranes.

We then tested protection from nuclease activity via DNase I digestion (Figure 2a). Following lipid treatment, 1.5 μg of each variant was incubated with 24 units of enzyme for 24 h at 37 °C, and total remaining DNA was quantified by PicoGreen fluorescence and compared to non-nuclease-treated controls. The 0 outer handle sample showed little protection with 30.4 ± 2.3% DNA remaining, whereas 84.6 ± 7.2% remained in the 48 outer handle sample, clearly

Figure 2. Bulk encapsulation yield and in vitro immune activation. (a) Encapsulation yield of outer handle DNO variants was estimated by PicoGreen dye membrane exclusion (red), and protection from nuclease was assayed with DNase I (blue). ELISA assay measurements of (b) IL-6 and (c) IL-12 cytokine production by splenocytes after incubation with N-DNO, E-DNO, and 50 nm vesicles for 16 h, as well as nonactivated controls. (d) Flow cytometry measurement of splenocyte mean fluorescence after incubation with Cy5-labeled N-DNO, E-DNO, and negative control. (e) Flow cytometry forward- (cell size) and side-scattering (granularity) properties of splenocytes was used to define two populations. Small, low granularity cells (1) were analyzed separately from large, high granularity cells (2). (f) Histogram of population (2) fluorescence after incubation with Cy5-labeled N-DNO (purple), E-DNO (blue), and negative controls (red). (*a,b: p < 0.05, ANOVA + Dunnet’s test vs control, error bars indicate SEM).
showing that the envelope provides protection against nuclease digestion.

Negative-stain TEM revealed a far more dramatic impact of handle number on ultrastructure (Supporting Information Figure 5). Whereas most 0 handle DNOs were not associated with vesicles, those with 12 or 24 outer handles had most of their surface covered by vesicles formed external to the nanostructure frame. Imaging of 48 handle DNO revealed nanostructures within apparently complete and tightly associated lipid bilayers, as shown in Figure 1c. The number and density of attached lipid-conjugated oligonucleotides is therefore a key design parameter for DNO encapsulation. Successful encapsulation was also dependent on the liposome formulation. Removal of PEG-DOPE resulted in much larger reconstituted vesicles and no micelle fusion. We observed that cholesterol (∼5%) can be used in place of PEG-DOPE to similarly control micelle fusion and to produce E-DNO (data not shown). Inclusion of both PEG (5%) and cholesterol (15%) caused stabilization of reconstituted vesicles at smaller diameters, suggesting an inhibition of fusion and trapping of a putative intermediate of the encapsulation process (Supporting Information Figure 6).

We next used density equilibrium centrifugation in an iodixanol gradient to separate 48 outer handle E-DNO from excess lipid vesicles, resulting in a highly enriched population (Supporting Information Figure 7). The final recovery yield of E-DNO relative to starting number of DNO used in encapsulation and the fraction of fully encapsulated nanostructures were determined by PicoGreen quantification and membrane exclusion to be ∼20 and 90%, respectively. This yield allowed us to prepare quantities of E-DNO suitable for in vitro and in vivo studies.

A significant innate inflammatory response would be a serious impediment to many biomedical applications of DNA nanotechnology. As a gauge of this, we incubated Cy-5-labeled N- and E-DNO, as well as empty 50 nm vesicles, with immune cells isolated from mouse spleens. Similar to previously reported results,17,18 N-DNO activated a potent inflammatory cytokine response comparable to that produced from exposure to bacterial or viral nucleic acids. Interleuken-6 (IL-6) and IL-12 were produced at 136 ± 10- and 99 ± 5-fold above nonactivated cells (Figure 2b,c). In contrast, E-DNO and 50 nm vesicles induced 2.1 ± 0.3- and 0.9 ± 0.3-fold increases, respectively, in IL-6 production versus controls. Similarly, IL-12 was undetectable after incubation with E-DNO and vesicles. Flow cytometry showed that mean fluorescence of splenocytes incubated with N-DNO was 111 ± 8-fold higher than with E-DNO, which was equivalent to negative control cells (Figure 2d). N-DNO uptake was concentrated in a subpopulation of large, granular cells (Figure 2e,f) in which 89.0 ± 3.2% displayed very bright Cy5 signal. This same subpopulation showed low mean fluorescence and few positive cells (8.0 ± 2.3%) after incubation with E-DNO, a result that was confirmed by confocal microscopy (Supporting Information Figure 8). This splenocyte assay shows that nanostructure activation of, and uptake by, immune cells can be almost fully attenuated by encapsulation in a PEGylated lipid membrane.

Encouraged by this, we next aimed to characterize the in vivo pharmacokinetics of E-DNO. AlexaFluor750-labeled oligonucleotides, N-DNO, and E-DNO were intravenously injected into anaesthetized mice to allow tracking via whole animal optical imaging for 120 min postinjection. Elimination half-life of the oligonucleotide control was estimated to be 38.0 ± 0.8 min, with signal rapidly accumulating in the bladder immediately after injection (Figure 3a,b). The half-life (49.5 ± 1.0 min) and clearance pattern of N-DNO were equivalent to that of the oligonucleotide. Because nanoparticles larger than 6 nm are size-excluded from renal clearance,30 these data indicate that the structural integrity of N-DNO becomes compromised immediately after injection. In contrast, E-DNO displayed an estimated half-life of 370 ± 38 min (Figure 3c), comparable to similarly formulated PEGylated liposomes,4 with little bladder accumulation. In comparison to N-DNO, encapsulation increased the elimination half-life and relative bioavailability by factors of 9 and 17, respectively.

The stability of DNA nanostructures and devices has previously been probed during exposure to lysate from sodium dodecyl sulfate-treated mammalian cells,31 and after direct injection into Caenorhabditis elegans,32 with little degradation observed. Comparison to our findings is confounded by the presence of nuclease-inhibiting surfactant, distinct nanostructure designs, and an alternative in vivo model system, yet the observed differences suggest that DNA-based nanomaterials may be sensitive to their environment in a design-dependent manner. A folate-targeted, DNA-based nanoparticle was also recently tested in a murine tumor model,33 displaying an elimination half-life (24.2 min) and kidney uptake similar to our non-encapsulated DNO, suggesting it may have suffered from a comparable, rapid degradation profile after injection.

Finally, we profiled biodistribution by imaging organs harvested 120 min postinjection (Figure 3d,e). No significant differences were observed between the distribution profiles of oligonucleotide and N-DNO on an organ-by-organ basis, and both displayed significant renal excretion with signal in urine of 86.3 ± 3.6 and 88.5 ± 2.9% (% of total, photons/g), respectively. In contrast, E-DNO urine accumulation reached only 11.0 ± 4.4% of AlexaFluor750 signal at this time point, while maintaining 85.0 ± 4.4% in blood. Dual labeling of the E-DNO membrane (rhodamine) and DNA nanostructure (AlexaFluor750) allowed us to compare biodistribution of both components through multiplex imaging (Supporting Information Figure 9).
No significant differences were observed between the two components, suggesting that they remained associated throughout imaging.

As a last comparison, we examined the biodistribution of 50 nm liposomes of the same formulation, hydrodynamic diameter, and concentration as E-DNO (Extended Data Figure 9). No significant differences in rhodamine fluorescence were observed between the major organs. Yet surprisingly, the 50 nm liposomes cleared more rapidly than E-DNO, with twice as much signal (38.0 (3.2% vs 20.1 (4.3%, respectively) measured in urine. This was contrasted with 58.0 (2.5% and 76.2 (3.3% signal remaining in blood of liposome and E-DNO injected animals. Although it requires more in-depth follow-up studies, it would be interesting if DNO acts as a stabilizing endoskeleton and reduces clearance of liposomes.

CONCLUSIONS
This virus-inspired E-DNO displays favorable in vitro and in vivo properties, in stark contrast to non-enveloped DNO which activates a potent immune response and displays rapid degradation after injection. Using this design strategy as a starting point, many different biomedical applications can be conceived through integration of additional functions. Because our E-DNO incorporates a contrast agent and has appropriate dimensions and pharmacokinetic properties, it could easily be adapted for tumor detection. Addition of ligands to the outer membrane leaflet would add specificity for target molecules or cells, and such ligands could potentially be spatially organized by the DNA "capsid shell" using transmembrane linkers. Many different triggers could be implemented to drive an internal molecular cascade or a mechanical reconfiguration of the nanostructure frame (e.g., photon-fuelled, thermally or chemically induced conformational switch, pH-sensitive lipids), which could be used for contrast agent unmasking, drug release, or more sophisticated behaviors that have been difficult to achieve using conventional nanomaterials or multicomponent nanodevices.

A more distant but highly exciting prospect is the development of autonomous devices having functions approaching that of virus particles (e.g., cell entry via attachment and fusion) or primitive immune cells (e.g., input-based therapeutic response), and which could provide significant advances in diagnostics and therapeutics.
between liquid N2 and a room temperature water bath, then at 850 rpm, room temperature for 1 h. The solution was then put on a dry N2 line, and the film was placed under vacuum overnight.

terol, PEG2K-PE, and rhodamine-PE (79.2, 15.0, 5.0, 0.8) in glucose to undetectable levels prior to DNA nanostructure purification.

Experiments were prepared in the same manner, except with a MgCl2 and SYBR Safe stain, in 0.5 M Tris (pH 8.0), 1 mM EDTA (1 × TE), and 10–22 mM MgCl2. The solution was then subjected to a thermal annealing ramp on a Tred 2 Petrit thermal cycler (Bio-Rad) according to the following schedule: 80 °C for 5 min, decrease to 65 °C at 5 min/°C, incubate at 65 °C for 20 min, decrease to 25 °C at 20 min/°C.

The products were separated in a 1.5% agarose gel with 10 mM MgCl2 and SYBR Safe stain, in 0.5 × TBE buffer + 10 mM MgCl2 for 3 h at 60 V. The leading bands were extracted and imaged by TEM. We opted for a 14 mM MgCl2 concentration for the synthesis of stock nanostructures.

Negative-Strain Transmission Electron Microscopy. TEM imaging was carried out by dropping 3.5 μL of product onto a plasma-treated carbon Formvar grid. This was incubated for 1 min, wicked away onto filter paper, 3.5 μL of 2% uranyl formate (in H2O, w/v) was added for 30 s, then wicked away. Imaging was carried out on a JEOL 14000 transmission electron microscope.

Purification of Synthesized Nanostructures. Solutions of folded DNA nanostructures were concentrated and then purified by glycerol gradient ultracentrifugation, as described elsewhere.

Liposomes Preparation. Liposomes were prepared using standard techniques. We produced a mixture of DOPE, PEG2K-PE, and rhodamine-PE (molar ratio 94.2, 5.0, 0.8), or DOPE, DOPS, PEG2K-PE, and rhodamine-PE (79.2, 15.0, 5.0, 0.8) in chloroform in a glass tube. Chloroform was evaporated using a dry N2 line, and the film was placed under vacuum overnight. We added a volume of encapsulation buffer (5 mM Tris-HCl, 1 mM EDTA, 10 mM MgCl2, 10 mM NaCl) to the film. Typically, we would add 300 μL of buffer to a total of 4.5 μmol of lipid. The film was resuspended by shaking on a Thermomixer (Eppendorf) at 850 rpm, room temperature for 1 h. The solution was then put through seven rounds of freezing and thawing by passing between liquid N2 and a room temperature water bath, then extruded with 21 passes through a Mini-Extruder (Avanti Polar Lipids) using a 0.2 μm polycarbonate membrane. Liposomes were stored at 4 °C, light-protected for up to 4 weeks.

The 50 nm empty vesicles prepared for in vitro and in vivo experiments were prepared in the same manner, except with a 50 nm extrusion membrane after extrusion through the 200 nm membrane.

Lipid – DNA Conjugate. The lipid–DNA conjugate was synthesized as described elsewhere.

DNA Nanostructure Encapsulation. In brief, DNA nanostructures were encapsulated by first annealing lipid–oligonucleotide and fluor–oligonucleotide conjugates to the nanostructure in a surfactant buffer. The annealed product was purified by glycerol gradient, then mixed with liposomes. This was then dialyzed to remove surfactant, and the product was purified, concentrated, dialyzed against a buffer appropriate to downstream experiments, and characterized. Sample volumes listed below were found to be scalable.

In a typical experiment, a 400 μL solution was prepared, containing 20–200 μg/mL DNA with 15% glycerol and 2% OG as surfactant. The solution was incubated for a minimum of 2 h at 35 °C on a Tred 2 Petrit Cycler (Bio-Rad).

The annealed product was purified from excess lipid–DNA and fluor–DNA conjugates by centrifugation. Glycerol gradients were prepared using solutions of 15% glycerol + 2% OG and 45% glycerol + 7% OG in an encapsulation buffer. The annealed product was placed on top of the gradients and centrifuged for 2.5 h at 41,000 rpm (for SW-41 rotor tubes). The gradients were then fractionated, and appropriate fractions were combined and concentrated back to the starting volume (i.e., 400 μL) using an Amicon 30K device.

The concentration of the final product was determined by UV absorbance at 260 nm on a Nanodrop system with disposable cuvettes. The volume was transferred into a 2.0 mL microcentrifuge tube. Liposomes were added by transferring a 0.5 × volume of prepared liposomes into the solution (i.e., 200 μL). This was mixed on the Thermomixer at 450 rpm, room temperature for 1 h. A volume of encapsulation buffer equivalent to the current total volume (i.e., 600 μL) was added and mixed gently. The entire solution was then transferred into an appropriately sized 7K MWCO Slide-a-Lyzer dialysis cassette (Thermo Scientific). The cassette was floated in 2 L of encapsulation buffer for 3 days for small samples (e.g., 120 μL) or 7 days for large volumes (e.g., 12 mL). Buffer was replaced every second day in all cases.

After dialysis, the sample was recovered from the dialysis cassette and concentrated using an Amicon column pre-treated with encapsulation buffer. Enveloped nanostructures were separated from excess lipids by equilibrium centrifugation using iodixanol (Optiprep reagent, Sigma-Aldrich). We prepared a working volume of 54% iodixanol from the stock 60% solution by mixing with 0.1 × volume of 10 × encapsulation buffer. We then mixed the 54% iodixanol/Encapsulation Buffer solution with 1 × Encapsulation Buffer to prepare equal volumes of 35, 28, 18, and 8% iodixanol/encapsulation buffer. In the case of the 35% step, the encapsulated DNA nanostructure sample was placed in place of 1 × encapsulation buffer to dilute the 54% iodixanol solution. The volumes were layered into ultracentrifuge tubes and centrifuged at maximum allowable speed for 16 h, 4 °C. The gradient was fractionated, and 50 μL of each fraction was transferred into a 96-well fluorescence plate (BD Bioscience) and imaged on the Typhoon system (GE Healthcare Life Sciences). The fractionation was profiled by quantifying fluorescent signals in each fraction using ImageJ (NIH). The images were background-subtracted, then a circular region-of-interest (ROI) was placed over each well, and integrated density was measured. The integrated density values were plotted against fraction number to profile component distribution. Appropriate fractions containing E-DNO (based on Cy5 or AlexaFluor750 signal) were concentrated and washed with encapsulation buffer using an Amicon centrifugation filter device or, in some cases, were first dialyzed against a buffer appropriate for downstream experiments (e.g., phosphate-buffered saline, Gibco).

The encapsulated, purified products were characterized by TEM, DLS, and fluorescence, and the encapsulation yield was determined by a PicoGreen assay (described below). DLS was carried out on a Nano ZS (Malvern) using standard settings.
Fluorescence analysis was carried out on a Fluorolog (Horiba) using the following settings. For rhodamine fluorescence of vesicles, excitation was at 550 nm and emission scanned from 575 to 700 nm. For Cy5 nanostructure fluorescence, excitation was at 625 nm and emission scanned from 650 to 800 nm. For AlexaFluor750 fluorescence, excitation was at 700 nm and emission scanned at 725–900 nm.

**PicoGreen Dye Exclusion Assay.** The PicoGreen stain exclusion assay differentiates between membrane-enclosed and non-enclosed DNA based on the stain’s inability to cross a lipid bilayer. Measurements are made in parallel in encapsulation buffer and buffer with OG that destabilizes the membrane and allows the stain access to total DNA content of a sample. The assay is carried out in a 384-well black fluorescence assay plate (Greiner).

A standard curve was first prepared from the DNA nanostructure stock solutions. First 200 μL of 5 μg/mL was prepared in encapsulation buffer, then 6× 1.2 dilutions were prepared from this. Next 100 μL of each unknown sample was prepared by dilution with encapsulation buffer (typically 1.5–1.50), aiming for the concentration of the diluted sample to be within the standard curve range. Two solutions of PicoGreen stain were prepared by diluting the stock solution 1:200 in either encapsulation buffer, or encapsulation buffer + 2% OG. For each standard or sample to be measured, 10 μL of both PicoGreen solutions was pipetted into 3× wells of the 384-well plate. Ten microliters of the standards and samples was then added to the 6× wells, and the plate was incubated 5 min and light-protected. Fluorescence of the PicoGreen stain was measured using a SpectraMax M5 plate reader (Molecular Devices) by excitation/emission at 480/520 nm.

For analysis, standard curves were plotted for median fluorescence values of both the OG-negative and OG-positive buffers. These were used to calculate the DNA concentration of unknown samples in the two buffers. The concentration of DNA in the OG-containing buffer equals “total DNA”, whereas that in the encapsulation buffer was “non-encapsulated DNA”. Encapsulation yield is then calculated by

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\text{Encapsulation yield = } \frac{(\text{total DNA} – \text{non-encapsulated DNA})}{\text{total DNA}} \times 100\%
\]

**Nuclease Protection Assay.** Sensitivity to nuclease activity was determined using DNase I (New England Biolabs). The encapsulation process was carried out on 0, 12, 24, and 48 outer handle DNA nanostructure variants at a point prior to purification by float-up centrifugation. One hundred microliters of each was transferred into PCR tubes (n = 3 for DNase I-positive and DNase I-negative of each variant, 12 tubes in total). Ten units of DNase I were added to 3× replicates of each variant, and an equivalent volume of DNase I buffer was added to 3× replicates as negative controls. All samples were incubated for 24 h at 37 °C on a Tetrat 2 Peltier thermocycler. Following this, the PicoGreen assay was used to determine the DNA remaining after nuclease digestion. The percent remaining was expressed as a ratio of the calculated DNA remaining in the DNase I-positive samples over the DNase I-negative samples for each design.

**Spleenocyte Activation Assay.** RPMI media (Gibco) used for splenocyte tissue culture were adjusted to maintain the stability of DNA nanostructures. MgCl₂ was added to 6 mM by addition of MgSO₄·7H₂O to maintain osmolarity. Cell viability and cytokine response in this medium was validated prior to data collection. To heat-inactivate nuclease activity present in FBS, serum was heat-treated for 30 min at 56 °C in 1 mL volumes on a Thermomixer at 1000 rpm. Heat-inactivated serum was added to RPMI at 10%. Penicillin-streptomycin (Gibco) was added at an appropriate dilution.

Spleens were obtained from female 8 week old C57Bl/6 mice (Charles River). Two spleens were processed for each experiment. They were transferred into 70 μm cell strainers (BD Falcon) and were dissociated using a sterile syringe plunger. The single cell suspension was washed through the cell strainer with 25 mL RPMI (Gibco) into 1 mL tubes. The suspension was transferred to a 50 mL Falcon tube, and PBS was added to a total volume of 40 mL. This was centrifuged at 500g for 5 min, and the supernatant was removed and discarded. PBS (40 mL) was added, the pellet was resuspended to wash the cells and dilute nuclease enzymes, and the sample was respun. A total of 3× washes were used to remove nuclease activity carried over from the tissue. Two milliliters of ACK lysis buffer (Lonza) was added, and the cells were gently resuspended and incubated for 7 min. Following this, 20 mL of PBS and 20 mL of RPMI media were added, and the suspension was centrifuged as above. The supernatant was discarded, and 15 mL of RPMI media was added. After sitting for several minutes to allow settling of large cell and tissue aggregates, the upper 10 mL of suspension was transferred to a new 15 mL Falcon tube.

Cell concentration was determined, and 1 × 10⁶ live cells were transferred in 450 μL to 4× wells of a 48-well plate for each sample to be assayed. Then, 50 μg/mL solutions of Cy5-labeled N- and Cy5/-rhodamine E-DNO were prepared (quantification via PicoGreen). A solution of 50 nm vesicles was prepared, with a concentration equivalent to E-DNO based on rhodamine absorbance. Fifty microliters of the nanostructure or vesicle samples was transferred into the wells (n = 4). These were incubated for 16 h in a CO₂ incubator at 37 °C. Supernatants were removed and centrifuged for 10 min at 500g, transferred to new tubes, and assayed for cytokine expression immediately or stored at −80 °C.

Cytokine concentrations were determined by enzyme-linked immunosorbent assays (R&D Systems) following the protocols exactly.

**Flow Cytometry.** Cells pelleted from the above splenocyte experiment were fixed by addition of 1.0 mL of 4% parafomaldehyde-hyde/PBS solution for 15 min. Cells were pelleted and washed 1× with PBS, then resuspended in 500 μL of PBS. Flow cytometry was carried out on a LSRFortessa (BD) using a 640 nm laser line with 670/30 nm emission filter to determine Cy5 fluorescence in samples (10000 events/measurement, n = 4). Gating was performed on splenocytes and “granulocytes” (large, granular cell population) by gating upon the FSC versus SSC dot plot. Identical gates were applied to all samples. After gating, a negative population was defined using the histogram obtained for media-only negative control samples. Samples that showed a rightward peak shift were determined as positive. Gates were applied to Cy5 histograms for splenocytes and granulocytes.

**Confocal Imaging.** Confocal imaging was carried out on splenocytes obtained from the above assay. The splenocytes were stained with a 20 mM Hoescht 33342 solution for 20 min, then resuspended in PBS. Cells were transferred into glass bottom dishes for imaging of Cy5 and Hoescht staining using standard filter sets and methods on an SPS xmp inverted confocal microscope (Leica). All images were acquired with a 63×1.2 water objective, 405 diode laser (Hoechst) and white light laser tuned to 575 nm (Alexa 568). Emission was collected with Leica Hybrid Detectors from 415 to 550 nm for Hoescht and 585–795 nm for Alexa 568. Optical zoom was used for higher magnification image.

**In Vivo Optical Imaging.** Optical imaging experiments were carried out on an IVIS Spectrum instrument (PerkinElmer). Athymic C57 nude mice (Charles River) were anesthetized under 3% isoflurane and were then injected via tail vein with a 100 μL bolus of AlexaFluor750-labeled (AF750) N-DNO, AF750/ rhodamine-labeled E-DNO at 50 nm, or rhodamine-labeled 50 nm liposomes of the same formulation used for encapsulation. They were immediately transferred to the imaging system, and data collection was initiated. Animals were maintained at 2% isoflurane throughout imaging. Fluorescence images for kinetic analysis were acquired by a 30 s excitation at 745 nm and emission at 800 nm, every 4 min for a total of 30 images over 120 min postinjection.

Following collection of kinetic data, mice were immediately euthanized and blood was collected by cardiac puncture and transferred into heparin collection tubes. Urine and organs were harvested and were imaged on the IVIS system for 2 s at 570/ 620 nm (rhodamine) or 30 s at 745/800 nm (AF750). Analysis was carried out in ImageJ (NIH). In all cases, images were background-subtracted using a 50 pixel rolling ball radius. Pharmacokinetic analysis was carried out using the Freehand Selection tool to draw a ROI around the complete head and...
torso of the animal to just below the front legs. The integrated density was measured in each image. The data set was normalized to the value at 8 min postinjection, as this showed peak fluorescence after diffusion of the agents throughout the vascular system. Organ measurements were obtained using the Freehand Selection tool to draw ROIs around their perimeter and measuring integrated density. Fifty microliter volumes of blood and urine were measured. "Organ distribution" was calculated for each organ as a percent of the total measured arbitrary fluorescent units, with urine normalized to a 200 μL estimated bladder volume and blood to 2500 μL estimated total blood volume. The contribution of organ fluorescence from blood volume was calculated and subtracted from total fluorescence.

Statistics. Student’s t test and ANOVA with posthoc Dunnett’s or Tukey’s tests were performed using an excel plug-in, inerror-STAT-a.v.3 by Mario H. Vargas (Instituto Nacional de Enfermedad Respiratorias, Mexico).

Animal Use. All animal studies were performed in accordance with NIH guidelines, under approval of Harvard University’s Institutional Animal Care and Use Committee.

Conflict of Interest: The authors declare the following competing financial interest(s): We filed a provisional patent on this technology.

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Supporting Information Available: Methods, additional data, and a nanostructure design schematic and oligonucleotide sequences are available as Supporting Information. This material is available free of charge via the Internet at http://pubs.acs.org.

REFERENCES AND NOTES


