Efficient Targeting of Fatty-Acid Modified Oligonucleotides to Live Cell Membranes through Stepwise Assembly

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Supporting Information

ABSTRACT: Lipid modifications provide efficient targeting of oligonucleotides to live cell membranes in a range of applications. Targeting efficiency is a function of the rate of lipid DNA insertion into the cell membrane and its persistence over time. Here we show that increasing lipid hydrophobicity increases membrane persistence, but decreases the rate of membrane insertion due to the formation of nonproductive aggregates in solution. To ameliorate this effect, we split the net hydrophobicity of the membrane anchor between two complementary oligonucleotides. When prehybridized in solution, doubly anchored molecules also aggregate due to their elevated hydrophobicity. However, when added sequentially to cells, aggregation does not occur so membrane insertion is efficient. Hybridization between the two strands locks the complexes at the cell surface by increasing net hydrophobicity, increasing their total concentration and lifetime, and dramatically improving their utility in a variety of biomedical applications.

INTRODUCTION

Lipid-modified oligonucleotides1−3 facilitate uptake of siRNA,4 target DNA nanostructures to lipid bilayers,5 program assembly of 3D microarrays,6,7 enable preparation of live single cell microarrays,8,9 and function as vaccine adjuvants and immunotherapeutics.10−12 These uses are predicated on rapid, efficient, and stable partitioning of these amphiphilic molecules from solution into live cell membranes.13 We recently reported an approach for incorporating dialkylglycerol modified oligonucleotides (DAG) into cell membranes.10 DAG is useful for targeting DNA to the membranes of most cell lines, but limits the ultimate efficiency of incorporation into the bilayer over time.

To improve the concentration of lipid-anchored oligonucleotides in cell membranes, we reasoned that increasing the dialkyl anchor hydrophobicity would increase its thermodynamic stability when inserted into cell membranes.9,14−16 Indeed, previous studies demonstrated that longer lipids are more stable than shorter lipids when reconstituted into synthetic lipid bilayers.3 However, we found that DAG incorporation into live cell membranes (as opposed to synthetic systems) was exquisitely sensitive to alkyl chain length. The addition of even two methylenes completely inhibited partitioning into cell membranes.9 We hypothesized that this was due to a competing self-aggregation reaction17,18 and thus sought an alternative means of introducing greater hydrophobicity to the lipid anchors without aggregation.

EXPERIMENTAL SECTION

Synthesis of Lipid-Modified Oligonucleotides. Hexadecanoic (Palmitic) acid, octadecanoic (Stearic) acid, icosanoic (Arachidic) acid, docosanoic (Behenic) acid, tetracosanoid (Lignoceric) acid, N,N-diisopropylhexadecylamine (DIPEA), N,N-diisopropylcholoramidite (DIPC), N,N-dimethylformamide (DMF), methylamine, ammonium hydroxide, and piperidine were obtained from Sigma-Aldrich. HPLC grade acetonitrile, triethylamine, acetic acid, and dichloromethane (DCM) were obtained from Fisher Scientific. Monomethoxytritylamine (hexyl-2-cyanoethyl)-(N,N-diisopropyl)-phosphoramidite (amine phosphoramidite), standard phosphoramidites, and DNA synthesis reagents were obtained from Azco Biotech. Controlled pore glass (CPG) support, 1-O-dimethoxytrityl-hexyl disulfide,1-[(2-cyanoethyl)-(N,N-diisopropyl)]-phosphoramidite, 10-O-[1-propyl-3-N-carbamoylcholestyryl]-triethyleneglycol-1,2-0-succinoyl-long chain alkyllamino-CPG (3′-cholesterol-TEG CPG), and 2-dimethoxytritylmethyl-6-fluoromethoxybenzylamino-hexane-1-succinoyl-long chain alkyllamino-CPG (3′-amino-modifier C7 CPG) synthesis columns

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were obtained from Glen Research. All materials were used as received from manufacturer.

Oligonucleotides were synthesized on an Applied Biosystems Expedite 8909 DNA synthesizer. Amino and cholesterol modified DNA strands were synthesized using amine and cholesterol phosphoramidites (100 mM), respectively, using a custom 15 min coupling protocol. For the polythymine regions of the anchor strands (ANCH), the capping step was omitted in order to maximize yield. After synthesis of 5′ amino-modified DNA, the MMT protecting group was deprotected manually on the synthesizer by priming alternately with deblock and dry acetonitrile three times and watching for yellow elution. To ensure complete deprotection of the MMT group, the 5′ solid supports were also resuspended in a solution of 20% acetic acid/80% water shaking for 1 h at room temperature. The solid support was subsequently washed repeatedly with DMF, DCM, and acetonitrile with acetonitrile as the final wash and then dried with a speedvac system. For the 3′ amino-modified CPG, a solution of 20% piperidine in dimethylformamide was prepared and used to deprotect the CPG support for 10 min at room temperature, followed by DCM and DMF washes with DCM as the final wash. This procedure was repeated twice more to ensure complete deprotection of the Fmoc protecting group prior to coupling to the fatty acid. Fatty acid conjugated oligonucleotides were synthesized by coupling the carboxylic acid moiety of the fatty acid to amino modified oligonucleotides with a 3′ or 5′ free amine while on the solid support. The solid support was transferred to an eppendorf tube and resuspended in a solution of dichloromethane containing 200 mM fatty acid, 400 mM DIPEA, and 200 mM DIPC. The eppendorf tubes were sealed with parafilm, crowned with a cap lock, and shaken overnight at room temperature. The next morning, they were washed with DCM and DMF repeatedly and then cleaved off the solid support.

Oligonucleotides were cleaved from solid support with a 1:1 mixture of ammonium hydroxide/40% methylamine (AMA) for 1 h at 65 °C with a cap lock followed by evaporation of AMA with a speedvac system. Cleaved oligonucleotides were filtered through 0.2 μm Ultrafree-MC Centrifugal Filter Units (Millipore) to remove any residual CPG support before HPLC purification. Fatty acid modified oligonucleotides were purified from unmodified oligonucleotides by reversed-phase high-performance liquid chromatography (HPLC) using an Agilent 1200 Series HPLC System equipped with a diode array detector (DAD) monitoring at 260 and 300 nm. Purifications used 100 mM triethylamine acetate (TEAA, pH 7) H2O/acetonitrile array detector (DAD) monitoring at 260 and 300 nm. Puriﬁed oligonucleotides were used for surface passivation. For the preannealing experiment, a 1 μM solution of C18/C16 and a 0.3 μM CHIR99021 and 1 μM PD0325901. The cells were lifted from their substrate using accutase and grown to a density of 2–3 x106 cells per cm2. MCF10A cells were kindly provided by Professor Jayanta Debnath (UCSF) and were cultured as previously described.1 Low passage, primary fibroblast, HMEC, and pre-adipocyte cell lines were provided by Jim Garbe (LBNL) and cultured in M87 media supplemented with chola toxin.

Surface Preparation for Cell Binding. Lyophilized 5′-amino-modiﬁed DNA was resuspended in a buffer of 60 mM sodium citrate, 450 mM sodium chloride, pH = 7.0. DNA was patterned onto PDMS using the spin coating apparatus described previously.2 Stocks of 250 μM ss-DNA strands were diluted to 1 μM with this buffer. The ss-DNA was then resuspended in distilled water and lyophilized again for 10 min at room temperature. This solution was used to resuspend the cell pellet after the final wash from media and gently agitated for an additional 10 min at room temperature. This was compared to normal labeling using these same strands at these same concentrations. All reported values are the average of three or more independent measurements, with error bars indicating standard deviations. Graphs were produced using the Prism software package (Graphpad). The heatmap was produced using the R software package (R), specifically using the ggplot2 library.

Measuring Aggregation by Dynamic Light Scattering (DLS). PBS CMF (UCSF CCF) was diluted by a 0.2 μm vacuum ﬁlter. Stock solutions of 250 μM ss-DNA strands were then diluted to 1 μM with this ﬁltered PBS prior to transfer to cuvette for measuring by DLS on a Wyatt Technology DynaPro Protein Solutions utilizing the DYNAMICS software package ver 6.10.1.2. Particle size was determined by cumulants analysis. All samples were prepared separately and measured in triplicate.

Surface Preparation for Cell Binding. Lyophilized 5′-amino-modiﬁed DNA was resuspended in a buffer of 60 mM sodium citrate, 450 mM sodium chloride, pH = 7.0. DNA was patterned onto aldehyde-silanized glass (Schott) using a micropipette (Figure 4c) or a Nano eNablar (BioForce Nanciences; Figure 4d,e). Slides were reduced with NaBH4 (Sigma) and passivated with both SigmaCote (Sigma) and Phuronic F108 acid before use. Cells labeled with lipid-DNA were allowed to settle onto patterned glass within a PDMS-based flow cell for 30 min. Flow cells were ﬂushed with ice cold PBS, and only cells hybridized to the surface via DNA were retained. Patterned cells were imaged using phase contrast settings and reconstructed using tiling algorithm (Zen Software, Zeiss). For experiments with mouse embryonic stem cells, no sigmatec was used for surface passivation.

Programmed Cell Assembly. For quantiﬁcation of programmed assembly efﬁciency, CellTracker Green CMFDA and CellTrace Far
Red DDAO-SE (Invitrogen) stocks were prepared to a concentration of 10 mM in anhydrous DMSO. Cells were resuspended in 10 μL stain in serum-free media for 30 min at 37 °C followed by 15 min in media supplemented with 10% fetal bovine serum before proceeding through the labeling steps described above. After washing away unreacted DNA, cells were resuspended at 1 × 10^6 cells/mL. Green cells were mixed with far-red cells at a ratio of 1:60 with 10^6 cells per 200 μL of ice cold PBS. Mixtures were then agitated at 150 rpm for 10 min in an Ultra-Low Attachment 24-well plate (Corning). This mixture was pelleted and resuspended in ice cold PBS before quantifying via flow cytometry or sorting via fluorescence activated cell sorting (FACS). The labels were reversed to ensure unbiased quantitation. For programmed assemblies of the embryonic stem and pancreatic beta islet cells, CellTracker Green CMFDA was used to stain the ESCs and the islet cells were left unstained. Cells were mixed with anchor strands 1 and 2, respectively, and assembled as described above. Clusters containing at least 1 green cell were purified from the unassembled cell population using a FACS Aria II (UCSF Laboratory for Cell Analysis).

**MCF10A Acinus Formation and Imaging.** MCF10A cells were cultured as described above. Cells were collected and stained with CellTracker Green CMFDA, as described above or left unstained. The green and unstained cells were subsequently labeled with anchor strands 1 and 2, respectively, and assembled as described above. Clusters containing at least 1 green cell were purified from the unassembled cell population using a FACS Aria II (UCSF Laboratory for Cell Analysis) and grown for 48 h in 3D on-top cultures in 8-well chamber slides (Lab-Tek) which were performed as previously described using growth-factor-reduced lE CM lots with protein concentrations between 9 and 11 mg/mL (Matrigel; BD Biosciences) (Debnath et al., 2003). After 48 h, the 3D cultures were fixed with 4% Paraformaldehyde in PBS. The 3D cultures were stained, as previously described by Debnath et al. (2003). Structures were stained with rat anti-human αν-integrin antibodies (Millipore clone NKL50H3-MAB1578) for the primary and Alexa-568 conjugated goat antirabbit antibodies (Invitrogen) for the secondary. Alexa-464 conjugated phalloldin (Invitrogen) and 1X DAPI in PBS was used to stain the actin cytoskeleton and nuclei, respectively. Confocal images were acquired on Zeiss Axio Observer Z1 equipped with a Yokogawa spinning disk unit and an EM-CCD camera.

## RESULTS AND DISCUSSION

Previous reports show that complementary cholesterol-bearing oligonucleotides can be stably targeted to liposomes and supported lipid bilayers via hybridization. We envisioned further increasing the hydrophobicity of the membrane anchors to further stabilize duplexes in live cells, rather than artificial lipid bilayers. To prevent aggregation of these more hydrophobic molecules, however, the two strands would need to be added sequentially to cells, rather than as a prehybridized duplex. Under conditions of stepwise addition, a first Anchor strand (Anch) partitions into the lipid bilayer but remains in rapid equilibrium with the medium. A second, co-Anchor (cA) strand is subsequently added and also establishes rapid equilibrium between the lipid bilayer and the medium. However, upon encountering the first strand through diffusion in the phospholipid bilayer, the two strands hybridize, increase the total hydrophobicity of the now doubly anchored duplex and, thus, slowing their exchange with the medium (Scheme 1).

To explore this strategy, we used fatty acid amides (FA) as more synthetically tractable membrane anchors than previously reported phospholipids or cholesterol. Fatty acids are widely commercially available and do not require chemical modification before coupling. Additionally, the conjugation reaction to DNA is not highly water sensitive and requires only one reverse phase purification step after coupling.

### Scheme 1. Stepwise Assembly of Fatty-Acid (FA)-Modified ssDNA into Cell Membranes

![Scheme 1](image-url)

(a) Structure of the Anchor (Anch) and co-Anchor (cA) strands incorporating a lock region for Anch/cA-strand annealing and a polythymine spacer region. A handle region provides adhesion to surfaces and a means of quantifying cell membrane incorporation using complementary fluorescently labeled probes. (b) Model for stepwise assembly of membrane anchored DNA dupplexes. FA-DNA molecules insert into the lipid bilayer but remain in rapid equilibrium with the surrounding cell medium. A second, complementary FA-modified oligonucleotide similarly establishes a rapid equilibrium with the cell membrane, but can also hybridize with the first strand in the membrane. The additional FA-anchor in the hybridized duplex alters the equilibrium, locking the co-anchored complex in the membrane. Elevated hydrophobicity can also trigger non-productive aggregate formation in solution.

Consistent with past studies, a single FA anchor does not stably label cell membranes when compared to DAG or doubly cholesterol-anchored DNA (Figure S1). For example, a 100 base Anch strand linked to stearic acid (C_{36}) via a 3′-amide (5′-Anch_{100}C_{16}) did not yield significant DNA incorporation after incubation with cells and washing (Figure 1, column 3). However, addition of a second, 20 base complementary coanchor (cA-) strand linked to palmitic acid (C_{16}) via a 3′-amide (3′-cA_{20}C_{16}) dramatically increased cell labeling to near that of the DAG and doubly anchored cholesterol (Figure 1, column 6 and Figure S1). No increase was seen upon addition of a noncomplementary 3′-cA_{20}C_{16} Strand (Figure 1, Column 5), indicating that at least two FA anchors, linked noncovalently through Watson–Crick base pairing in the “lock” region (Scheme 1a), are necessary for stable incorporation.

We found that the number of base pairs in the lock region correlated with initial labeling and retention of oligonucleotides over time, both at 0 and 37 °C. This effect saturated between 15 and 20 bases (Figure S2). Labeling was dose-dependent and occurred without altering cell viability over the examined range of 0.5 to 5 μM (Figure S2). Encouragingly, even these unoptimized molecules were capable of programming cell–cell and cell–surface adhesion of model cell lines with results comparable to DAG when incorporating 60 base polythymine spacers (Figure S3).

These initial findings suggested we could achieve additional improvements in cell membrane incorporation by increasing the length and thus hydrophobicity of FA anchors. We therefore synthesized a series of 5′-Anch_{100} strands conjugated to saturated FAs between 16 and 24 carbons in length. These Anch strands behaved as predicted when added stepwise to cells in concert with 3′-cA_{20}C_{16}. Increased lipophilicity of FAs enhanced the labeling efficiency and showed substantial

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improvement over both DAG and cholesterol linked oligonucleotides (Figure 2A). Anch strands with enhanced hydrophobicity also demonstrated improved retention over time at physiological temperature (Figure S4). Unlike the 5′-Anch100-C18 and 3′-cA20-C16 combination, stepwise addition of more hydrophobic Anch and cA strands was essential for preventing competing aggregation reactions (Figure 2B). Prehybridizing 5′-Anch100-C24 and 3′-cA20-C16 strands led to dramatically reduced cell membrane incorporation compared to prehybridized 5′-Anch100-C18 and 3′-cA20-C16 strands (Figure 2B). Together, these results support the notion that splitting the hydrophobicity of dual-anchored species across two complementary oligonucleotides added stepwise to cells prevents aggregation and improves labeling.

These data suggested we could achieve further increases in cell labeling and stability by increasing the hydrophobicity of the cA strand in addition to the Anch strand. Surprisingly, increasing the hydrophobicity of the cA strand anchors did not yield additional gains in cell labeling. For instance, stepwise addition of 5′-Anch100-C24 and 3′-cA20-C24, which maximizes hydrophobicity for both strands, actually decreased DNA incorporation when compared to 5′-Anch100-C24/3′-cA20-C16. We investigated this effect by assaying a panel of molecules in which the fatty acid on the Anch and cA strands was varied systematically and independently. We found that increasing hydrophobicity specifically on the coanchor strand decreased labeling (Figure 2C). Indeed, 5′-Anch100-C24/3′-cA20-C16 inserted far more efficiently into cell membranes than 5′-Anch100-C16/3′-cA20-C24 despite containing identical number of phosphodiester bonds and methylene groups.

To explain this trend, we hypothesized that the ratio of anchor hydrophobicity to oligonucleotide length (and thus charge) determines the extent of aggregation. If this were the case, short oligonucleotides would be more prone to aggregation than equivalently modified longer oligonucleotides. To test this notion, we used dynamic light scattering (DLS) to examine the relationship between FA anchor length, oligonucleotide length, and relative aggregation. Both scattered light intensity (Figure 3A) and particle size (Figure S5) correlated with the length of the FA conjugated to the cA strand. In contrast, very little light scattering was observed for any FA conjugated to the 100 base anchor strand (Figure 3B). These results suggested that adding additional bases to the cA strand, increasing its net size and charge, would destabilize aggregates through Coulombic or steric repulsion while simultaneously allowing for increased hydrophobicity of its FA anchor. We therefore synthesized a series of 3′-cA20-C24 strands incorporating an additional 10, 20, or 30 bases at the 5′ end. Consistent with our expectations, DLS revealed an inverse

Figure 1. Anchor (Anch) and complementary co-Anchor (cA) strands together enhance ssDNA targeting and retention in cell membranes. Fluorescence was measured with flow cytometry and normalized to a C16/C18 DAG-ssDNA control. cA-MM is a 20 base coanchor strand with a DNA sequence noncomplementary to the Anch strand. Error bars are standard deviation of at least three independent measurements.

Figure 2. Lipid hydrophobicity affects cell labeling efficiency of Anch, cA, and prehybridized strands. (a) Membrane incorporation compared to DAG for duplexes with 3′-cA20-C16 strands and Anch strands bearing FA anchors of increasing length. (b) Membrane incorporation of 3′-cA20-C16 and different Anch strands when added stepwise (white bars) or after preannealing (black bars). (c) Heatmap relating average membrane labeling (n = 3) to combinations of Anch and cA strand FA anchor lengths. Error bars are standard deviation of at least three independent measurements.

Figure 3. Ratio of FA to oligonucleotide length determines the extent of aggregation. (a) Light scattering from solutions of 3′-cA20 strands as a function of FA anchor length. (b) Light scattering from solutions of 3′-cA-C24 anchored oligonucleotides as a function of the number of DNA bases. The red box indicates the same strand, 3′-cA50-C24. (c) Membrane incorporation compared to DAG for combinations of 3′-cA-C24 with increasing numbers of DNA bases. (d) Incorporation vs time for DAG and 5′-Anch100-C16/3′-cA20-C24-DNA. Error bars are standard deviation of at least three independent measurements.
relationship between the number of bases and aggregation (Figure 3B). Moreover, the best of these molecules, 3’-cA26-C24, increased cell labeling in combination with 5’-Anch100-C24/3’-cA26-C16 to nearly 7-fold of DAG (Figure 3C). This combination of molecules also showed a dramatic increase in lifetime at the cell surface compared to DAG (Figure 3D). We calculated that the initial rate of decay of these fully optimized strands from the cell surface was nearly 100-fold lower than DAG.

Given the improved cell labeling enabled by stepwise addition of C24 conjugated oligonucleotides, we investigated whether they could be used for programming cell—cell and cell—surface adhesion for cell types that were inaccessible with DAG. Using even unoptimized 5’-Anch100-C24/3’-cA26-C16 we found efficient adhesion between ESCs and complementary 7 μm diameter spots of DNA patterned on glass surfaces, allowing the preparation of live, single cell microarrays with more than 95% occupancy. By comparison, DAG yielded only 40% binding using identical conditions (Figure 4A). We also used stepwise assembly of FA-anchored DNA to prepare 3D microtissues using stepwise assembly of membrane anchored adhesive oligonucleotides. (a) Single cell microarrays prepared with murine ESCs labeled with Anch100-C24/cA26-C16 or DAG. (b) MCF10A clusters assembled around 1 cell tracker green-stained cell (arrow) establish polarity after 48 h in Matrigel (scale bar is 20 μm). (c) Representative cluster of sorted ESC/mouse cells (scale bar is 10 μm) and labeling efficiency of β cell islet cells. Error bars are standard deviation of at least three independent measurements.

**CONCLUSION**

In conclusion, stepwise assembly of membrane-anchored oligonucleotides is a modular strategy for targeting DNA to cell membranes with improved efficiency and stability. Insertion of oligonucleotide duplexes into membranes occurs via two FA-anchors with higher net lipophilicity compared to previously reported anchors. Competing self-aggregation is prevented by separating the dual anchors between two molecules that are added sequentially to cells, as well as by balancing the ratio of hydrophobicity to oligonucleotide length. This strategy facilitates new applications such as DNA-mediated adhesion in primary cells, murine ESCs, and pancreatic β cells, cell types that show little to no labeling with DAG. An additional benefit of these molecules is their streamlined synthesis compared to previous methods. We anticipate that the structure/function relationships defined here will prove useful in other applications utilizing lipid-modified oligonucleotides or amphiphiles including vaccine adjuvants, siRNA delivery, and structural DNA nanotechnology.

**ASSOCIATED CONTENT**

Supporting Information

Oligonucleotide sequences, MALDI data, and additional figures. This material is available free of charge via the Internet at http://pubs.acs.org.

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Notes

The authors declare no competing financial interest.

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