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**Aptamer photoregulation in vivo**

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Aptamer photoregulation in vivo

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Contributed by Robert Langer, October 21, 2014 (sent for review July 31, 2014)

The in vivo application of aptamers as therapeutics could be improved by enhancing target-specific accumulation while minimizing off-target uptake. We designed a light-triggered system that permits spatiotemporal regulation of aptamer activity in vitro and in vivo. Cell binding by the aptamer was prevented by hybridizing the aptamer to a photo-labile complementary oligonucleotide. Upon irradiation at the tumor site, the aptamer was liberated, leading to prolonged intratumoral retention. The relative distribution of the aptamer to the liver and kidney was also significantly decreased, compared to that of the free aptamer.

Aptamers are single-stranded nucleic acids that have emerged as a promising class of therapeutics owing to their relative ease of synthesis and high affinity and selectivity toward a range of targets including small molecules, proteins, viral particles, and living cells (1–6). Aptamers can fold into well-defined conformations and are more resistant to enzymatic degradation than other oligonucleotides (7–9). Aptamers have been suggested for imaging applications because their relatively small size and molecular mass (∼10 kDa) allow fast tissue penetration and clearance from blood (10, 11). The same characteristics make aptamers promising for effective delivery of diagnostic and therapeutic agents to tissues or organs. However, nonspecific accumulation of aptamers in normal tissues is undesirable (12–15) because it diminishes the proportion of aptamer that targets the desired tissue. This can adversely affect the therapeutic index of the aptamer; this may be particularly true if the aptamer is conjugated to a drug or drug delivery device. Moreover, aptamers themselves can have nonspecific toxic effects (16, 17).

Aptamers can be either generated by systematic evolution of ligands by exponential enrichment (SELEX) (18–20), or spatiotemporal regulation of aptamer activity in vivo has not been achieved, whereby activity would be enhanced in target tissues and not others. Here, we report a strategy to provide light-triggered control of aptamer function and distribution in vivo.

Light is an excellent means of providing external spatiotemporal control of biological systems (21–26). Many strategies have been developed to incorporate photosensitive groups in oligonucleotides that can control cellular function or affect biological pathways or gene expression by light (24–29). Of particular interest, such approaches can be used to provide spatiotemporal control of gene activation (24). Here we hypothesized that light triggering can be used to achieve spatiotemporal control of binding of an aptamer injected systemically to its target tissue in vivo, which would have implications for control of delivery of therapeutic aptamers and/or conjugated drugs or drug delivery systems. We designed a photo-triggerable system whereby the aptamer of interest is inactivated by hybridization to a photo-labile complementary oligonucleotide. Upon irradiation, the complementary sequence breaks down, releasing the functional aptamer (Fig. 1). The aptamer of interest is the single-stranded DNA 26-mer aptamer AS1411 (A\textsubscript{1411}; sequence: 5′-GGT GGT GGT GGT TGT GGT GGT GG-3′) that binds with high affinity and selectivity to nucleolin (30–32), which is overexpressed on the cell membrane of several types of cancer cells, including the 4T1 breast cancer cells used here (33–35). A\textsubscript{1411} has been used for cancer targeting in vitro and in vivo (35, 36). A complementary photo-triggerable inhibitory oligonucleotide (OliP) was designed [sequence: 5′-CCCA CCA/CCA CAA/CA A C-3′, where // indicates photo-labile 1-(2-nitrophenyl)ethyl bonds (37); Scheme S1].

Results

We examined the formation and light-triggered dehybridization of double-stranded A\textsubscript{1411}/OliP in vitro using Förster resonance energy transfer (FRET). A\textsubscript{1411} was labeled with a Cy3 fluorophore at the 3′ end (Cy3-A\textsubscript{1411}) and OliP was modified with a quencher of Cy3 (Iowa Black RQ) at the 5′ end (Q-OliP); the change of FRET signal between the fluorophore and quencher would allow evaluation of the photoactivation process (Fig. 2 A and B). The addition of Q-OliP to Cy3-A\textsubscript{1411} in solution quenched the fluorescence of Cy3 due to FRET (Fig. 2 C), confirming the formation of the A\textsubscript{1411}/OliP hybrid structure. Irradiation (365 nm light at 5 mW/cm\textsuperscript{2}) of the Cy3-A\textsubscript{1411}/Q-OliP hybrid resulted in an increase in fluorescence intensity at the emission wavelength of Cy3 (Fig. 2 D), indicating release of the Cy3-A\textsubscript{1411} strand from the hybrid, which peaked in 5 min (Fig. S1). There was no change in fluorescence intensity upon irradiating Cy3-A\textsubscript{1411} hybridized to a quencher-bearing DNA strand with the same sequence as Q-OliP but without photo-cleavable bonds (Q-Oli; Fig. S2), suggesting that breakage of the photo-labile bonds by light was crucial to the release of the aptamer from the hybrid. Native polyacrylamide gel electrophoresis was performed to further confirm the light-triggered dehybridization of A\textsubscript{1411}/OliP; ~95% of the OliP was cleaved after 5 min of irradiation (365 nm light at 5 mW/cm\textsuperscript{2}; Fig. S3).

The ability of A\textsubscript{1411}/OliP hybrid to target cancer cells was investigated in vitro. Fluorescent Cy3-A\textsubscript{1411}/OliP was incubated with 4T1 breast cancer cells (nucleolin-positive) at 37 °C for 2 h. Fluorescence microscopy revealed minimal uptake into cells exposed to 0.5 μM Cy3-A\textsubscript{1411}/OliP compared with cells incubated with free Cy3-A\textsubscript{1411} (Fig. 3 D), indicating that hybridization to OliP inhibited aptamer activity in vivo has not been achieved, whereby activity would be enhanced in target tissues and not others. Here, we report a strategy to provide light-triggered control of aptamer function and distribution in vivo.

Significance

The importance of this research is in the demonstration of the spatiotemporal regulation of aptamer activity in vitro and in vivo that can be achieved by a photo-triggered strategy. The aptamer’s cell-binding capability was blocked by hybridization with a photo-cleavable complementary oligonucleotide. Irradiation at the tumor site liberated the aptamer from the duplex, leading to prolonged intratumoral retention. Furthermore, the relative distribution of the aptamer to the tumor was enhanced in relation to uptake by the liver and kidneys. Our strategy may provide an approach to improving the therapeutic indices of aptamer-based medicines.

Author contributions: L.L., R.T., and D.S.K. designed research; L.L., R.T., H.C., and W.W. performed research; L.L., R.T., R.L., and D.S.K. analyzed data; and L.L., R.T., R.L., and D.S.K. wrote the paper.

The authors declare no conflict of interest.

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the activity of A1411. Irradiation of Cy3-A1411/OliP (365 nm, 5 min, 5 mW/cm²) induced cell uptake comparable to that seen with free Cy3-A1411, suggesting that irradiation released functional A1411 from the hybrid. Quantification by flow cytometry revealed that the fluorescence intensity of 4T1 cells incubated with Cy3-A1411/OliP was approximately three times less than that of cells treated with Cy3-A1411 (∼P < 0.005; n = 4). Upon irradiation (365 nm light at 5 mW/cm² for 5 min), the fluorescence intensity of cells incubated with Cy3-A1411/OliP increased ∼2.5-fold (∼P < 0.005, n = 4), that is, to a level similar to that of cells treated with free A1411 (Fig. S3B). In contrast, cellular uptake of Cy3-A1411/OliP without photocleavable bonds did not increase upon irradiation (Fig. S4), supporting the view that it was the photocleavable nature of OliP that enabled light-responsive cell binding by A1411/OliP.

Intratumoral photoactivation of A1411/OliP was demonstrated by intravital microscopy (38, 39) in nude mice with 4T1 tumors (diameters ∼6–8 mm) on their mammary fat pads. Mice were administered A1411 or A1411/OliP (250 nmol/kg i.v., both labeled with the fluorescent dye Cy3. FITC-dextran (molecular mass ∼70 kDa) was also injected i.v. to highlight the vasculature (40). Injection of free Cy3-A1411 resulted in strong extravasated fluorescence in the tumor tissue 10 min after injection. Administration of Cy3-A1411/OliP resulted in minimal fluorescence in the same time frame (Fig. 4). Irradiation of the tumor site (365 nm light at 200 mW/cm² for 3 min; see SI Materials and Methods, section S6 for the rationales for irradiation timing and Figs. S5 and S6) immediately after injection of Cy3-A1411/OliP resulted in more extravasated fluorescence throughout the tumor tissue compared with that without light irradiation. These results suggest that A1411/OliP could be dehybridized to liberate A1411 upon irradiation at the tumor site in vivo.

Fluorescence imaging of tumor sections 2 h postadministration showed distribution of Cy5-A1411 throughout the tumor tissue (Fig. 5), whereas A1411/OliP was mainly localized around blood vessels, presumably because of inhibition of the aptamer’s binding to cells. [The 2-h time point was selected because pilot studies suggested that free (unbound) A1411/OliP was still present in tumors in large quantity at earlier time points; see also Fig. 6B.] When the tumor site was irradiated for 3 min immediately after administration of A1411/OliP, Cy5 fluorescence was found to be distributed throughout the tumor at necropsy 2 h later.

To examine the effect of light triggering on biodistribution, nude mice bearing s.c. 4T1 tumors were administered Cy5-labeled A1411 or A1411/OliP i.v. then underwent quantitative whole-body fluorescence imaging (Fig. 6). The tumors were placed at the neck so that signal from the liver and other organs with significant nonspecific uptake would not interfere with signal from the tumor. To
minimize autofluorescence, the mice were fed a purified diet for 1 wk before experiments (41). Fluorescence from A1411 was readily detected in tumors 2 h after injection (Fig. 6A) but not in animals injected with A1411/OliP. Irradiation at the tumor site (365 nm light at 200 mW/cm² for 3 min) immediately after i.v. injection of A1411/OliP resulted in strong fluorescence at the tumor site, similar to that in the group receiving A1411, and 1.4-fold higher than that in mice receiving A1411/OliP without irradiation (Fig. 6B). The rate of clearance from tumor of A1411/OliP after irradiation (t1/2 = 1,124.8 s; Fig. 6B) was slower than that of A1411/OliP (t1/2 = 977.2 s). Of note, A1411/OliP showed similar intratumoral fluorescence intensity 2 h after administration with and without irradiation (200 mW/cm², 3 min) (Fig. S7), suggesting that increased binding of A1411/OliP was due to photo-cleavage of OliP and not a direct effect of irradiation on tissue, such as enhanced capillary permeability. These results suggest that A1411 was liberated from the hybrid upon irradiation, rapidly taken up by tumors (Fig. 4), and was retained there. Examination of irradiated H&E-stained sections of tissues (tumors and overlying skin) did not show signs of tissue injury compared with nonirradiated controls (Fig. S8).

Quantitative fluorescence imaging of tumors 2 h after injection showed that the intratumoral fluorescence signal intensity (median radiance) in the light-triggered A1411/OliP group was similar to that in the group treated with free A1411 (P > 0.1, n = 5, Fig. 6C and D) and was 2.2-fold higher than that in the group that received A1411/OliP without irradiation (P < 0.05, n = 5, Fig. 6C). Irradiation also increased aptamer accumulation in tumor relative to levels in other organs. The median ratio of liver to tumor radiance in the group that received A1411/OliP with light triggering was 2.5-fold less than that in the group treated with A1411 (P < 0.05).

**Discussion**

We have demonstrated a strategy for regulating aptamer activity and biodistribution by light triggering. In our design, hybridization with a photo-cleavable complementary oligonucleotide (A1411/OliP) blocked the aptamer’s tumor-targeting capability, which was restored when the aptamer was released by photo-triggering. Hybridization also changed the aptamer’s biodistribution, possibly by two phenomena: (i) enhanced binding to tumor upon irradiation in relation to nonspecific tissue binding, and (ii) A1411 [and some other aptamers (7)] has a configuration that protects it from nuclelease digestion (42) and is reported to have very different biodistribution from other DNA molecules that undergo rapid organ uptake followed by rapid degradation and renal clearance of breakdown products (10, 43, 44). The stability of A1411 is greatly diminished by hybridization in A1411/OliP (Fig. S9).

Aptamers can be covalently bound to a wide variety of therapeutic agents, including chemotherapeutics or enzymes (45, 46), siRNA (47, 48), or drug-loaded nanocarriers (36, 49), to provide targeted drug delivery. The spatiotemporal control of drug delivery afforded by our approach may enhance the efficacy and therapeutic index of many drugs. Our strategy involves less synthetic workup than approaches that change oligonucleotide biodistribution via chemical modification (e.g., PEGylation) (14, 50–52).

Although UVA light (320–400 nm) has been widely used for photochemical regulation of biological activities in vitro and in vivo (21–26), UVA light possesses clinical limitations such as limited tissue penetration. Clinical applicability could be enhanced by use of two-photon technology whereby near-infrared light, which can penetrate tissues more deeply, could be used to cleave two-photon photolabile groups (26, 53). Fiberoptic endoscopy could also be used to use UVA light deep within the body (54, 55). Brief irradiation with 365 nm light at low energy is not considered a risk for skin cancer (56, 57).

**Materials and Methods**

A full description of materials and methods is provided in SI Materials and Methods.

**A1411/OliP Synthesis.** A1411/OliP stock solution was prepared by adding OliP to a solution (PBS, 150 mM NaCl, pH 7.4) of DNA aptamers labeled with a dye with a final ratio of A1411/OliP of 1:1. The solution was annealed and stored at 4 °C overnight to allow full hybridization.

![Fig. 4. Representative intravital microscopic images of orthotopic 4T1 tumors 10 min after i.v. administration of Cy3-labeled A1411 or A1411/OliP, without or with light triggering at the tumor site (365 nm light at 200 mW/cm² for 3 min) immediately after injection. Green, Cy3-labeled A1411 or A1411/OliP; red, FITC-dextran (70 kDa) that demarcates blood vessels; blue, intratumoral collagen matrix; yellow, the localization of A1411 or A1411/OliP in blood vessels. The accumulation of A1411 and A1411/OliP in extravascular spaces (indicated by white dashed ovals) with light triggering was observed. Two animals were imaged in each group.](image-url)

![Fig. 5. Intratumoral aptamer distribution. (A) Representative fluorescence images of intratumoral distribution of Cy5-labeled A1411 and A1411/OliP (red) 2 h after i.v. injection, without or with irradiation (365 nm at 200 mW/cm² for 3 min) immediately postinjection. Blue, cell nucleus; green, antibody against CD31 staining blood vessel endothelial cells; yellow, colocalization of red and green. (B) Quantitation of the data in A. Numeric data are medians ± quartiles; n = 4. Asterisk indicates P < 0.01. ns, P > 0.05.](image-url)
Whitney (1 mL), they were fixed with 4% (wt/wt) formaldehyde for 10 min at room temperature. After incubation for 2 h, the tumors are indicated by red dashed circles. (B) Time course of intratumoral Cy5 fluorescence over 2 h, derived from integration of the photoradiance (photons per second per square centimeter per steradian [p/s/cm²]) in A. n = 5. (C) Intratumor accumulation of Cy5 2 h after injection. Data are medians ± quartiles. Asterisk indicates P < 0.05. (D) Representative fluorescence images of organs and tumors 2 h after i.v. injection. (E) The ratio of photoradiance in livers to that in tumors and kidneys to tumors for mice that received A₁₄₁₁ or A₁₄₁₁/OliP with irradiation. Asterisks indicate P < 0.01 (n = 5 for all groups). Numeric data are medians ± quartiles.

**Cellular Uptake Analysis.** 4T1 cells were grown in 12-well plates in RPMI-1640 medium, supplemented with 100 units/mL aqueous penicillin G, 100 mg/mL streptomycin, and 10% (w/vt) FBS (all from Life Technologies) at concentrations to allow 70% confluence in 24 h (i.e., 2 × 10⁵ cells/cm²). On the day of experimentation, the medium was replaced with Opti-MEM medium (500 mL) containing 0.5 μM Cy3-labeled A₁₄₁₁ or A₁₄₁₁/OliP probes with or without light triggering (365 nm light at 5 mW/cm² for 5 min). After incubation for 2 h, the cells were washed with PBS (2 × 500 μL per well) and treated with 0.25% trypsin with EDTA for 5 min (Life Technologies). After the cells were washed with PBS (2 × 1 mL), they were fixed with 4% (w/vt) formaldehyde for 10 min at room temperature, washed with PBS (2 × 1 mL) and stored in 1 mL PBS with 1% (w/vt) BSA solution at 4 °C for flow cytometry analysis.

**The 4T1 Tumor Model and in Vivo Imaging.** Immunodeficient 6- to 8-wk-old nu/nu nude mice were purchased from Charles River Laboratories and maintained under pathogen-free conditions for all animal studies. The study protocol was reviewed and approved by the Massachusetts Institute of Technology Committee on Animal Care. For subcutaneous 4T1 tumor models, 4T1 cells were injected in the dorsal aspect of the neck with 1 × 10⁶ cells/0.1 mL in HBSS were injected into mouse mammary fat pads. Tumor length and width were measured with calipers, and the tumor volume was calculated using the following equation: tumor volume = length × width / 2. Mice whose tumors reached ∼100–200 mm³ were used in subsequent experiments. The mice were injected intravenously (i.v.) with Cy5-labeled A₁₄₁₁ or A₁₄₁₁/OliP at an aptamer dose of 250 nmol/kg. In light-triggering experiments, the tumor site was illuminated by UV light for 3 min (365 nm, 200 mW/cm²) immediately after injection. Whole body fluorescence imaging was performed with an IVIS imaging system (IVIS spectrum, Caliper Life Sciences) with excitation and emission wavelengths of 640 and 680 nm, respectively.

**Statistical Analysis.** Data which were reasonably normally distributed were described with means and standard deviations and compared with t tests. Otherwise, data were presented as median ± quartiles and differences between groups were assessed with a Mann–Whitney U test. All data analyses were performed using Origin 8 software.

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