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Photo-reactive oligodeoxynucleotide-embedded nanovesicles (PROsomes) with switchable stability for efficient cellular uptake and gene knockdown†

Beob Soo Kim, 📭 a Mitsuru Naito, 📭 Rimpei Kamegawa, a Hyun Jin Kim, 📭 Ryo lizuka,^c Takashi Funatsu,^c Shingo Ueno,^d Takanori Ichiki,^{ad} Akihiro Kishimura^e and Kanjiro Miyata **

A photo-responsive nanovesicle is fabricated by polyion complex (PIC) formation between poly(ethylene glycol) (PEG)-block-polypeptides and photo-reactive oligodeoxynucleotides (PROs)/anti-sense oligonucleotides (ASOs). The ultraviolet (UV) light triggers reversible crosslinking between PROs and ASOs in the vesicular membrane, providing the nanovesicle with switchable stability under physiological conditions. The resulting nanovesicle allows efficient cellular internalization, leading to significant UV-triggered gene knockdown in cultured cells.

Polymeric vesicles, termed polymersomes, have been widely investigated as a desirable platform for biochemical and biomedical applications because they can stably encapsulate both hydrophobic and hydrophilic substances in the membrane and hollow aqueous cavity, respectively.1 Indeed, various drugs, such as anticancer drugs, proteins/enzymes, and genetic materials, have been loaded into polymersomes for controlled drug release and targeted drug delivery.2 Notably, such drug payloads can be selectively released from polymersomes by controlling the membrane permeability with various stimuli. ^{2b,3} As an external physical stimulus, light is one of the attractive stimuli because it can trigger drug release through precise spatial and temporal control without additional substances.⁴

Very recently, we reported nano-sized vesicular PICs that consist of a PIC membrane embedded with small nucleic acids,

i.e., small interfering RNAs (siRNAs), and PEG-block-polypeptides.⁵ In this formulation, siRNAs played two roles; one was a structural component for nanovesicle formation and the other was a pharmaceutical substance. This siRNA-embedded nanovesicle demanded additional stabilization to maintain the vesicle structure under physiological conditions. To this end, glutaraldehyde was employed as a crosslinker between polypeptide segments (or primary amines) but not siRNAs to avoid ineffective release of siRNA payloads. However, glutaraldehyde, which is a homobifunctional crosslinking reagent, likely elicits uncontrollable and irreversible reactions, including self-polymerization and nanoparticle aggregation. Thus, more controllable and reversible crosslinking techniques are required for diverse applications of small nucleic acid-embedded nanovesicles.

Previously, PROs were synthesized by attaching photoreactive vinyl groups, e.g., cyanovinylcarbazole (CNVK) and ^{CNV}K-modified D-threoninol (^{CNV}D), to the nucleobases. The PROs installed with CNVK or CNVD can be reversibly reacted with an adjacent pyrimidine base in the complementary strand via [2+2] cycloaddition under UV irradiation at 366 nm (UV₃₆₆) for crosslinking and 312 nm (UV₃₁₂) for de-crosslinking. These PROs with high reactivity, selectivity, and reversibility have been utilized for in situ DNA manipulation, cytosine to uracil conversion, gene repair, and genetic engineering.^{6,7}

This UV-triggered reversible crosslinking chemistry has inspired us to fabricate a PRO-embedded vesicular PIC, termed PROsome, with switchable stability that enables the stable loading of ASOs in extracellular milieu and the triggered release of ASOs in cells for efficient gene knockdown (Fig. 1). The PROsome is fabricated through PIC formation among three components, PROs as an anionic reversible crosslinker, ASOs as an anionic therapeutic agent, and PEG-block-polypeptides as a cationic membrane component. In this way, UV365-triggered crosslinking should stably hold ASO payloads in the PIC membrane, while UV312-triggered de-crosslinking can compromise the stability of PIC membrane for the facilitated payload release. Herein, we demonstrate the successful fabrication of

^a Department of Materials Engineering, Graduate School of Engineering, The University of Tokyo, 7-3-1 Hongo, Bunkyo-ku, Tokyo 113-8656, Japan. E-mail: miyata@bmw.t.u-tokyo.ac.jp

^b Center for Disease Biology and Integrative Medicine, Graduate School of Medicine, The University of Tokyo, 7-3-1 Hongo, Bunkyo-ku, Tokyo 113-0033, Japan

^c Graduate School of Pharmaceutical Sciences, The University of Tokyo, 7-3-1 Hongo, Bunkyo-ku, Tokyo 113-0033, Japan

^d Innovation Center of NanoMedicine, Kawasaki Institute of Inductrial Promotion, 3-25-14 Tonomachi, Kawasaki-ku, Kawasaki 210-0821, Japan

^e Center for Molecular Systems, Kyushu University, 744 Moto-oka, Nishi-ku, Fukuoka 819-0395, Japan

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Fig. 1 Schematic illustration for preparation and delivery strategy of PROsome.

PROsomes and their photo-responsive functionalities for enhanced gene knockdown in cultured cells.

First, vesicular PIC formation was verified for the mixture of PRO, ASO, and PEG-block-polypeptide in an aqueous solution. The structural information of oligodeoxynucleotides is summarized in Table S1 (ESI†). The PRO was designed to have 5 ^{CNV}D-installed adenine (A) units in 20-mer polyA for random crosslinking with pyrimidine, i.e., thymine (T) and cytosine, in ASO. The nuclearenriched abundant transcript 2 (NEAT2)-targeting ASO (asNEAT2) was selected to evaluate the gene knockdown potential because NEAT2 long non-coding RNA (lncRNA) plays as an oncogene transcript and is highly expressed in many types of human cancers, including human lung cancers.8 As a PEG-blockpolypeptide, PEG-block-poly[N-(5-aminopentyl)- α , β -aspartamide] (PEG-P(Asp-AP)) was synthesized to have a M_n of PEG: 2000 and a degree of polymerization of P(Asp-AP): 39, as confirmed by size exclusion chromatography (Fig. S1, ESI†) and ¹H NMR (Fig. S2, ESI†). Of note, this polymer composition can satisfy the criteria of a f_{PEG} (defined by 100 × [total weight of PEG in a PIC]/[total weight of components in a PIC]) of less than 10% for vesicular PIC formation under a charge-stoichiometric condition (defined by [number of amino groups in PEG-P(Asp-AP)]/[number of phosphate groups in ASO/PRO] = 1).9 Indeed, PEG-P(Asp-AP) has formed vesicular PICs with various polyanions, including siRNA, in previous studies.^{5,10} Here, PIC samples were prepared by vortex mixing of PEG-P(Asp-AP)/PRO/asNEAT2 at a molar charge ratio of 2/1/1 (f_{PEG} : 8%), followed by irradiation with UV₃₆₅ (1920 mJ cm⁻²).6 The formation of vesicular PICs was characterized by dynamic light scattering (DLS) and transmission electron microscopy (TEM). Regardless of UV₃₆₅ irradiation, PIC samples had a hydrodynamic diameter of approximately 80 nm with a narrow size distribution (polydispersity index (PDI): ~ 0.1) (Fig. 2a). The TEM image of UV₃₆₅-irradiated PICs clearly displays vesicular nanostructures with darker pool and brighter rim (Fig. 2b), similar to those of previously reported vesicular PICs. 10 Of note, the vesicular nanostructures were maintained after UV₃₁₂ irradiation (Fig. S3, ESI†), indicating that the UV₃₁₂ irradiation did not induce structural changes in vesicular PICs.

Next, the stability of PROsomes was investigated to confirm whether the ^{CNV}D moieties in PRO were effectively

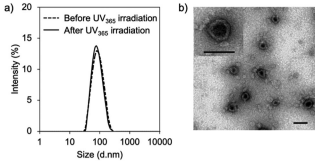


Fig. 2 Characterization of PICs prepared from PRO, asNEAT2, and PEG–P(Asp-AP). (a) Size distribution histograms obtained by DLS. (b) TEM image of PICs after irradiation with UV $_{365}$ (1920 mJ cm $^{-2}$). Inset: High magnification. Scale bars are 100 nm.

photo-crosslinked with asNEAT2 in the PIC membrane. The stability was examined by static light scattering (SLS) and DLS in terms of changes in scattered light intensity (SLI), size, and PDI of PROsomes under a physiological salt condition. UV₃₆₅untreated PROsomes, i.e., non-crosslinked (NX-)PROsomes, instantly reduced relative SLI (Fig. 3a). In contrast, the PROsomes treated with 1 min UV365 irradiation, i.e., crosslinked (X-)PROsomes, exhibited apparently constant SLI even after 48 h incubation. Also, the DLS results revealed that X-PROsomes maintained their initial size and PDI without significant changes in physiological milieu (Fig. 3b). These results indicate that the UV365 irradiation to PROsomes dramatically inhibited their structural changes in physiological milieu. Of note, considerable changes in relative SLI, size, and PDI of PROsomes were observed after treatment with less than 1 min UV365 irradiation (Fig. S4, ESI†), suggesting that 1 min UV365 irradiation was required for

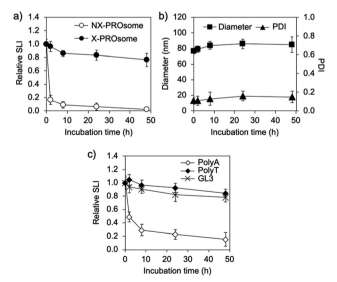


Fig. 3 Stability of PROsomes. Time-dependent changes in (a) relative SLI and (b) hydrodynamic diameter and PDI of PROsomes (PRO/asNEAT2) treated with UV $_{365}$ in 150 mM NaCl. (c) Time-dependent changes in relative SLI of PROsomes prepared from PRO/polyA, PRO/polyT, or PRO/asGL3 after UV $_{365}$ irradiation in 150 mM NaCl. All results are expressed as mean \pm SD (n = 4).

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the stabilization of nanovesicle structures under the physiological milieu.

To validate the effect of photo-crosslinking of PRO (or ^{CNV}D) with T in ASO, control PROsomes were further prepared from a series of oligodeoxynucleotide mixtures, i.e., PRO/polyA, PRO/polyT, and PRO/GL3 luciferase-targeted ASO (asGL3) (Table S1, ESI†), and then, were treated with UV₃₆₅ in the same manner as the PROsomes prepared from PRO/asNEAT2. PolyT and asGL3 including T units clearly suppressed the decrease in relative SLI, whereas such a suppression effect was not observed for polyA (Fig. 3c). This result clearly indicates that the photo-crosslinking between PROs (or CNVD moieties) and T-containing oligodeoxynucleotides should be essential for the significant stability of PROsomes in the physiological salt solution. It should be noted that both asNEAT2 and asGL3 containing six T units at different positions in 21-mer clearly improved the PROsome stability, similar to polyT, suggesting that 6 T units in ASO should be enough for the photo-reaction with PRO having five CNVD moieties in the PIC membrane.

The cyclobutane structure formed by the [2+2] cycloaddition between CNVD and T in oligodeoxynucleotides can be decrosslinked by photo-splitting reaction at UV₃₁₂ irradiation.^{6b} This de-crosslinking reaction makes PROsomes structurally unstable in the physiological environment. To confirm the reversibility of PROsomes, the time-dependent changes in SLI and hydrodynamic diameter of X-PROsomes in 150 mM NaCl solution were monitored under UV312 irradiation with a 15 W transilluminator (Fig. 4a and Fig. S5, ESI†). The relative SLI value of X-PROsomes rapidly decreased by only 0.5 min irradiation, associated with the considerable decrease in size of X-PROsomes. This result clearly demonstrates that the photosplitting reaction of ^{CNV}D occurred by UV₃₁₂ irradiation. The stability of X-PROsomes was additionally evaluated in fetal bovine serum (FBS)-containing media by fluorescence correlation spectroscopy (FCS) in terms of time-dependent changes in the hydrodynamic diameters of Alexa Fluor® 647-labeled ASO (AF647-ASO) (Fig. 4b). In this assay, two PROsome samples, X-PROsomes before and after UV₃₁₂ irradiation for 0.5 min, were prepared and incubated with 10% FBS. The hydrodynamic

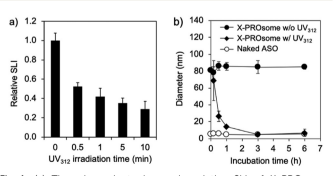


Fig. 4 (a) Time-dependent change in relative SLI of X-PROsomes (PRO/asNEAT2) under UV₃₁₂ irradiation in 150 mM NaCl solution. (b) Timedependent changes in hydrodynamic diameters of naked AF647-ASO and AF647-ASO-loaded X-PROsomes in the presence of 10% FBS before and after UV₃₁₂ irradiation, determined by FCS. All results are expressed as mean \pm SD (n = 3).

diameter of X-PROsomes without UV312 irradiation was kept approximately 90 nm, indicating that AF647-ASO was stably confined in the nanovesicles even in the serum-containing medium. In contrast, the hydrodynamic diameter of X-PROsomes was reduced immediately after 0.5 min UV312 irradiation and reached the similar size to naked ASO, suggesting that the nanovesicles were dissociated to release free AF647-ASO. These results demonstrate the photo-switchable stability of PROsomes.

The feasibility of PROsomes as an ASO carrier was then investigated using cultured human lung cancer, A549 cells. The cellular uptake efficiency of PROsomes was evaluated by flow cytometry (Fig. 5a). Naked AF647-asNEAT2 exhibited a modest cellular uptake efficiency, presumably because chemically modified ASOs with phosphorothioate (PS)-backbone and locked nucleic acid (LNA)-wings (i.e., PS-gapmers) have significant tolerability against nucleases. 11 The fluorescence intensity of cells treated with NX-PROsomes was comparable to that treated with naked AF647-ASO, suggesting that NX-PROsomes were unstable in the serum-containing medium and dissociated to free AF647-ASO. In contrast, X-PROsomes elicited a significantly enhanced cellular uptake efficiency, probably due to their higher tolerability in the serum-containing media. This enhanced cellular uptake of X-PROsomes can be explained by some reasons; (i) charge neutralization of ASO and (ii) larger surface area (curvature) of vesicular PICs compared with naked ASO, for facilitated adsorptive endocytosis. 12 The additional uptake inhibition assays suggest that X-PROsomes were mainly uptaken by endocytic pathways, including macropinocytosis, because the uptake amount of X-PROsomes was significantly decreased by incubation at 4 °C and treatment with cytochalasin D (Fig. S6, ESI†).

Next, the intracellular ASO release from X-PROsomes was observed by confocal laser scanning microscopy (CLSM). In this experiment, UV312 was directly irradiated to the cells to destabilize X-PROsomes through de-crosslinking. Thus, the UV₃₁₂ irradiation condition was first optimized to avoid the UVinduced cytotoxicity and DNA damage. 13 The UV312-treated cells

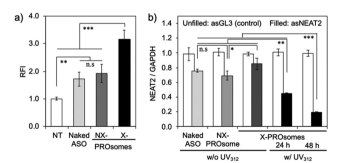


Fig. 5 (a) Cellular uptake efficiency of naked AF647-asNEAT2 and AF647asNEAT2-loaded PROsomes with or without UV₃₆₅ irradiation, determined by flow cytometry. The A549 cells were incubated with each sample at 100 nM AF647-ASO for 6 h. Relative fluorescence intensity (RFI) was calculated by normalizing to those from non-treated cells. (b) Gene knockdown efficiency of PROsomes determined by qRT-PCR. The A549 cells were incubated with naked ASOs or PROsomes at 100 nM ASO. All results are expressed as mean \pm SD (n = 4, *p < 0.05, **p < 0.01, and ***p < 0.001).

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showed a negligible effect on their viability in the case of an irradiation time within 10 min (Fig. S7, ESI†). Also, the concentration of messenger RNA (mRNA) harvested from the cells was determined as an indicator of DNA damage after treatment with UV312 at different irradiation times (Fig. S8, ESI†). No significant reduction in mRNA concentration was observed for the cells treated with UV₃₁₂ irradiation within 1 min. Thus, the UV₃₁₂ irradiation for 0.5 min was selected for the intracellular ASO release assay. The X-PROsomes were prepared from AF647-ASOs and AF546-ASOs at a molar ratio of 1:1 and incubated with cultured A549 cells for 24 h before UV₃₁₂ irradiation. After UV₃₁₂ irradiation, the cells were further incubated for 24 or 48 h to track the intracellular ASO release. The obtained CLSM images display that the yellow spots that result from the colocalization of AF647-ASOs and AF546-ASOs were reduced by UV₃₁₂ irradiation (Fig. S9a, ESI†), which is more quantitatively shown by the calculated Pearson's correlation coefficients (Fig. S9b, ESI†). These results indicate that the UV₃₁₂ irradiation triggered the time-dependent ASO release from X-PROsomes through de-crosslinking in the cells.

Eventually, the gene knockdown efficiency of PROsomes in cultured A549 cells was evaluated by quantitative reversetranscription polymerase chain reaction (qRT-PCR). We designed the gene knockdown experiment as illustrated in Fig. S10, ESI,† where UV₃₁₂ was irradiated for 0.5 min at 24 h post-transfection to induce de-crosslinking of X-PROsomes inside cells. Consistent with the cellular uptake study, the similar modest gene knockdown efficiencies ($\sim 30\%$) were observed for naked asNEAT2 and asNEAT2-loaded NX-PROsomes (Fig. 5b). Despite the high cellular uptake efficiency, X-PROsomes without UV312 irradiation elicited only 10% decrease in NEAT2 level, suggesting inefficient ASO release from X-PROsomes presumably due to the covalent crosslinking between PRO and ASO. In contrast, the gene knockdown efficiency of X-PROsomes was significantly enhanced up to $\sim 55\%$ and $\sim 80\%$ in 24 h and 48 h post-incubation, respectively, by 0.5 min UV₃₁₂ irradiation. Importantly, the X-PROsomes equipped with asGL3 as a control sequence induced almost no reduction in NEAT2 level. These results demonstrate the UV-triggered gene knockdown ability of X-PROsomes in an ASO sequence-specific manner. Of note, the X-PROsomes exhibited negligible cytotoxicity even at a high concentration of ASO (800 nM asGL3) (Fig. S11, ESI†).

In summary, the photo-responsive nanovesicle was successfully fabricated using PRO. The PROsomes were composed of anionic oligodeoxynucleotide mixtures and cationic block copolymers through their electrostatic interactions. While the UV365-crosslinking between CNVD-installed PRO and T-containing ASO provided the PROsomes with the structural durability in physiological milieu, the UV₃₁₂-de-crosslinking destabilized the PROsomes directed toward the ASO release inside cells, resulting in the enhanced gene knockdown. This is the first study demonstrating the stimuli-responsive vesicular PIC for enhanced functionalities. The photo-reversible strategy can provide versatile applications of multifunctional polymeric nanovesicles for codelivery of various macromolecular drugs

with their controlled release profiles. In the future study, we will further investigate the stability of the PIC membrane according to the number of photo-reactive moieties in the PRO strand. Also, other hydrophilic species will be simultaneously encapsulated into the hollow aqueous cavity of PROsomes for their cooperative functionalities.

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Conflicts of interest

There are no conflicts to declare.

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