A High-Throughput Platform for Formulating and Screening Multifunctional Nanoparticles Capable of Simultaneous Delivery of Genes and Transcription Factors

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Abstract: Simultaneous delivery of multiple genes and proteins (e.g., transcription factors; TFs) is an emerging issue surrounding therapeutic research due to their ability to regulate cellular circuity. Current gene and protein delivery strategies, however, are based on slow batch synthesis, which is ineffective, poorly controlled, and incapable of simultaneous delivery of both genes and proteins with synergistic functions. Consequently, advances in this field have been limited to in vitro studies. Here, by integrating microfluidic technologies with a supramolecular synthetic strategy, we present a high-throughput platform for formulating and screening multifunctional supramolecular nanoparticles (MFSNPs) self-assembled from a collection of functional modules to achieve simultaneous delivery of one gene and TF with unprecedented efficiency both in vitro and in vivo. We envision that this new approach could open a new avenue for immuno-therapy, stem cell reprogramming, and other therapeutic applications.

Research is currently directed toward transcription factors for the purpose of investigating disease therapy because of their ability to control cell behavior.[1–3] Transcription factors (TFs) are proteins that contain DNA-binding domains to recognize matching DNA sequences adjacent to the genes they regulate. Researchers have attempted to control transcription factors’ function by delivering the related DNA[15] or the transcription factor itself[2,3] into the cell to steer the cell’s fate towards a desired direction (e.g., apoptosis or cell differentiation). The transportation of transcription factors to a target location is especially challenging because most are easily decomposed by various enzymes in the body. Therefore, researchers have developed a wide variety of techniques for the delivery of genes and proteins.[4–12] However, the conventional approaches, including viral vectors,[4] nanocapsules,[5] nanoparticles,[6,7] liposomes,[7–9] copolymers,[10,11] and polypeptides[12] employed for optimization of delivery systems require empirical and multiple optimization cycles to repeat design/synthesis/assay processes. Given the enormous complexity of the biological system, it is unlikely that such a time- and cost-consuming developmental pipeline could lead to a crucial breakthrough in the search for optimal delivery vectors. As a result, most research has been limited to in vitro systems, apart from the recently reported multifunctional oligonucleotides.[15] Their advances in an in vivo system were highlighted[14] but still suffer from limitations on fast screening of the optimal combination because of their complex chemical synthesis. We proposed a convenient, flexible, and modular supramolecular synthetic strategy for delivery of intact (unmodified) TFs in a highly efficient manner.[15] Yet, continuous efforts need to be devoted to achieving in vivo applications.

We have combined integrated microfluidic technologies[16–24] with a supramolecular synthetic strategy[15,25–29] and report herein a high-throughput approach for formulating and screening multifunctional supramolecular nanoparticles (MFSNPs) that are capable of simultaneous delivery of gene and functional proteins with superb efficiency and control.
lable stoichiometry among individual payloads (Figure 1). As illustrated in Figure 1a, this approach utilized two microfluidic systems including a digital droplet generator (DDG; see Section S1.2–S1.5 and video in the Supporting Information) and a microfluidic cell array (Section S1.8) which were made by encapsulating the protein within a thin layer of degradable polymer of which the surface was linked with Ad groups; 2) the gene module can be any DNA of interest (i.e., pEGFP-encode DNA plasmid, pEGFP); 3) the function modules are a series of poly(ethylene glycol) (PEG) derivatives of which one end-group is terminated with an Ad group and the other end is terminated with a methyl group (Ad-PEG) or a functional peptide, such as RGD (Ad-PEG-RGD) (Section S1.10) and TAT (Ad-PEG-TAT) (Section S1.11); and 4) the scaffold module is CD-linked poly(ethyleneimine) (CD-PEI) (Section S1.12). This strategy can be used to produce generally applicable protein modules whose assembly properties are not affected by the core protein. Driven by the supramolecular synthetic strategy, genes and proteins can be effectively incorporated together to form MFSNPs grafted with various functional molecules (Ad-PEG, Ad-PEG-RGD, and Ad-PEG-TAT) on their surface to achieve desired structural stability, delivery specificity, and cell transfection capability, respectively. Moreover, because the gene and protein payloads are encapsulated inside the self-assembled MFSNPs, potential degradation during the delivery process can be avoided.

By modulating the inlet of the functional module, three categories of MFSNPs with different surface ligands were synthesized (Supporting Information, Figure S1). The 1st category of MFSNPs was decorated with only Ad-PEG; the 2nd category (RGD-MFSNPs) contained Ad-PEG and Ad-PEG-RGD; and the 3rd category (TAT-MFSNPs) bears Ad-PEG, Ad-PEG-RGD, and Ad-PEG-TAT with pre-identified ratios (Table S1). Each category contains 125 formulations of MFSNPs that were prepared by systematically altering the concentration of CD-PEI, BSA-Cy5, and HRP-RhB (Section S1.4). As a result, 375 formulations of MFSNPs were prepared automatically with 1 h (Table S2). After 30 min incubation, each MFSNP droplet (0.2 μL) was then automatically diluted with Opti-MEM media to 2 μL, and introduced into an individually addressed cell culture chip. After incubating the MFSNPs-treated cells in the cell array chip at 37 °C (5% CO2) for 24 h, the transfection and transduction efficiencies of each MFSNP formulation were quantified by fluorescence mi...

Figure 1. a) Two microfluidic systems, i.e., a digital droplet generator and a microfluidic cell array chip, were employed for formulating and screening of MFSNPs. b) Supramolecular synthetic strategy allows generation of a combinatorial library of gene- and protein-encapsulated MFSNPs by systematically altering the mixing ratio among the four functional modules, including i) proteins, ii) genes, iii) ligands (Ad-PEG, Ad-PEG-RGD, and Ad-PEG-TAT), and iv) a scaffold (CD-PEI).
croscopy-based single-cell image cytometry. A 3D plot (Figure 2a) summarizes the transfection and transduction efficiencies of the MFSNPs library, and shows that an optimal transfection/transduction performance was achieved using RGD/TAT-MFSNPs (3rd category) with a specific composition, that is, CD-PEI = 20 ng, Ad-PEG = 100 ng, Ad-PEG-RGD = 5 ng, Ad-PEG-TAT = 9 ng, pEGFP = 10 ng, BSA-Cy5 = 39 ng, and HRP-RhB = 27 ng in 200-nL PBS. Figure 2b shows the fluorescence images of cells treated with optimal RGD/TAT-MFSNPs, thus indicating the successful transfection of pEGFP and the successful transduction of both of the two proteins.

To gain insight on how the structural property of RGD/TAT-MFSNPs affects the observed co-delivery performance, the optimal synthetic parameters (Figure 2a,b; red *) were chosen for scale-up preparation (200 droplets, 40 μL). The RGD/TAT-MFSNPs are of spherical morphology with diameters of 100 ± 4 nm and surface charges of 7.2 ± 2.5 mV, which were confirmed by transmission electron microscopy (TEM) and dynamic light scattering measurements (DLS; Figure S5–S7). Furthermore, a colorimetric assay was employed for quantifying the enzymatic activity of RGD/TAT-MFSNP-encapsulated HRP intracellular by using o-dianisidine (3,3′-dimethoxybenzidine) as chromogenic substrates (Figure 2c). The time-dependent in vitro transfection and transduction performance of 100 nm RGD/TAT-MFSNPs was monitored in parallel by quantifying fluorescent signals associated with EGFP expression and colorimetric readouts correlated to HRP activity, respectively. As shown in Figure 2d, the highest co-delivery performance was achieved 24 to 48 h after RGD/TAT-MFSNP-treatment. Moreover, the other two formulations that led to suboptimal delivery performances of RGD/TAT-MFSNPs were also carefully examined (Figure S5 and Table S1). TEM images indicated that the size of those two suboptimal RGD/TAT-MFSNPs were larger than 200 nm (Figure S5d), which is unfavorable for high-performance delivery. The stability of the RGD/TAT-MFSNPs in various conditions was also confirmed by a series of studies (Figure S8–S14). In addition, no toxic effect was observed with RGD/TAT-MFSNPs (Figure S15).

The uniqueness of the high-efficient co-delivery vector[15] lies in the fact that it can simultaneously introduce a functionally complementary protein and gene that will lead to synergistic outcomes for regulating cellular circuitry. To exploit the use of RGD/TAT-MFSNP vector for delivery of functionally complementary gene and TF, pG5E4T-Fluc (a plasmid vector that contains GAL4-VP16 matching recognition sequences and a luciferase reporter), and GAL4-VP16 (a transcription factor fusion protein) were chosen for conducting in vitro and in vivo studies (Figure 3a). Again, GAL4-VP16 was first coated with a degradable polymer of which the surface was linked with Ad groups. After formulation and intracellular delivery of pG5E4T-Fluc and GAL4-VP16, the luciferase reporter in pG5E4T-Fluc was specifically activated by GAL4-VP16, generating a real-time readout (bioluminescence), thus reflect-
In vitro and in vivo co-delivery of functionally complementary gene and protein. a) Simultaneous delivery of gene (pG5E4T-Fluc) and functionally complementary protein (GAL4-VP16) with optimal 100 nm RGD/TAT-MFSNPs. b) Bioluminescence imaging and c) time-dependent luciferase expression of the 100 nm RGD/TAT-MFSNP-treated cells along with the control experiments based on pG5E4T-Fluc plasmid and GAL4-VP16. Error bars were obtained from three independent experiments. d) Monitoring in vivo synergetic effects of the co-delivered gene and protein after intravenous injection of 100 nm RGD/TAT-MFSNPs (containing 1 µg pG5E4T-Fluc and 6.6 µg Ad-n-GAL4-VP16) into nu/nu nude mice (n = 3) via the tail veins. e) Ex vivo experiments of 100 nm RGD/TAT-MFSNPs-treated mice were performed after 168 h post-injection.}

Figure 3. In vitro and in vivo co-delivery of functionally complementary gene and protein. a) Simultaneous delivery of gene (pG5E4T-Fluc) and functionally complementary protein (GAL4-VP16) with optimal 100 nm RGD/TAT-MFSNPs. b) Bioluminescence imaging and c) time-dependent luciferase expression of the 100 nm RGD/TAT-MFSNPs-treated cells along with the control experiments based on pG5E4T-Fluc plasmid and GAL4-VP16. Error bars were obtained from three independent experiments. d) Monitoring in vivo synergetic effects of the co-delivered gene and protein after intravenous injection of 100 nm RGD/TAT-MFSNPs (containing 1 µg pG5E4T-Fluc and 6.6 µg Ad-n-GAL4-VP16) into nu/nu nude mice (n = 3) via the tail veins. e) Ex vivo experiments of 100 nm RGD/TAT-MFSNPs-treated mice were performed after 168 h post-injection.

ing a synergistic outcome of the co-delivered payloads. By using the previously optimized synthetic parameters, 10 droplets (2 µl in total volume) containing desired RGD/TAT-MFSNPs were generated from CD-PEI (200 ng), Ad-PEG (1000 ng), Ad-PEG-RGD (50 ng), Ad-PEG-TAT (90 ng), pG5E4T-Fluc (100 ng), and GAL4-VP16 (660 ng) in the DDG. The resulting RGD/TAT-MFSNPs were introduced into a 96-well plate (containing NIH 3T3 cells, ca. 1 x 10^4 cells/well) along with control systems, including pG5E4T-Fluc plasmid and the protein module containing GAL4-VP16. After incubating (5% CO2) at 37°C for 24, 48, and 72 h, the cells were lysed for quantification of bioluminescence using either a plate reader or a cooled charge-coupled device (CCD) camera (IVIS, Xenogen) (Figure 3b,c). Compared to background-level bioluminescence intensities observed from the control experiments, the bioluminescence intensity of the RGD/TAT-MFSNP-based vectors was significantly higher, thus suggesting that the RGD/TAT-MFSNPs are capable of delivering both pG5E4T-Fluc and GAL4-VP16 simultaneously into the cells in a highly efficient manner as well as allowing both the protein and gene functioning synergistically.

Finally, the in vivo experiments were performed by injection of the RGD/TAT-MFSNPs that encapsulated pG5E4T-Fluc (1 µg) and GAL4-VP16 (6.6 µg) into nu/nu nude mice (n = 3) via the tail veins. As illustrated in Figure 3d and Figure S16, a strong bioluminescent signal was observed from the liver of the treated animal, thus indicating efficient delivery of the RGD/TAT-MFSNPs into the hepatocytes. The intensity of the signal at 168 h post injection decreased twofold in magnitude compared to that at 48 h post injection (Figures S16 and S17). Ex vivo results were consistent with bioluminescent imaging data (Figure 3c).

In conclusion, by integrating microfluidic technologies with a supramolecular synthetic strategy, we have demonstrated a new high-throughput approach for formulating and screening MFSNPs. Using our DDG, hundreds of formulations with variable delivery performance were programmed into the MFSNP combinatorial library. Subsequently, in vitro screening of the MFSNPs library in cell array chips identified specific formulations that lead to desired delivery performance. The optimal formulations were then employed to encapsulate functionally complementary TF and gene into MFSNP-based vectors, and the respective synergistic outcomes were achieved both in vitro and in vivo. This approach could open up a new avenue[10] for immunotherapy, stem cell reprogramming, and other therapeutic applications.

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