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# A Rapid Pathway Toward a Superb Gene Delivery System: Programming Structural and Functional Diversity into a Supramolecular Nanoparticle Library

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#### **Abstract**

Nanoparticles are regarded as promising transfection reagents for effective and safe delivery of nucleic acids into specific type of cells or tissues providing an alternative manipulation/therapy strategy to viral gene delivery. However, the current process of searching novel delivery materials is limited due to conventional low-throughput and time-consuming multistep synthetic approaches. Additionally, conventional approaches are frequently accompanied with unpredictability and continual optimization refinements, impeding flexible generation of material diversity creating a major obstacle to achieving high transfection performance. Here we have demonstrated a rapid developmental pathway toward highly efficient gene delivery systems by leveraging the powers of a supramolecular synthetic approach and a custom-designed digital microreactor. Using the digital microreactor, broad structural/functional diversity can be programmed into a library of DNA-encapsulated supramolecular nanoparticles (DNACSNPs) by systematically altering the mixing ratios of molecular building blocks and a DNA plasmid. In vitro transfection studies with DNACSNPs library identified the DNACSNPs with the highest gene transfection efficiency, which can be attributed to cooperative effects of structures and surface chemistry of DNACSNPs. We envision such a rapid developmental pathway can be adopted for generating nanoparticle-based vectors for delivery of a variety of loads.

<sup>&</sup>lt;sup>‡</sup>These authors contributed equally to the work.

# Keywords

supramolecular nanoparticle; gene delivery; digital microreactor; combinaltorial library; cyclodextrin; molecular recognition

## Introduction

Gene delivery constitutes one of the most critical steps in gene manipulation and therapy. <sup>1, 2</sup> By mimicking the size and function of viral vectors, numerous non-viral gene delivery systems based on biocompatible nanostructured materials, <sup>3–5</sup> *e.g.*, inorganic nanoparticles, <sup>6–9</sup> carbon nanotubes, <sup>10</sup> liposomes, <sup>11</sup> cationic polymers <sup>12</sup>, <sup>13</sup> and dendrimers, <sup>14</sup> have been developed to provide an alternative approach to the problems in viral gene delivery. However, an emerging challenge of nanoparticle-based delivery reagents is their low transfection performance. <sup>3</sup> In order to overcome this problem researchers have attempted to modulate <sup>15</sup> the structural (sizes <sup>16</sup> and shapes) and functional (surfaces chemistry and charges) properties of nanoparticles, <sup>15</sup> often requiring multiple optimization cycles to gradually improve the performance of nanoparticle-based delivery reagents. Such procedures have proven to be time-intensive and have demonstrated limited diversity over the past decades. Alternatively, it is believed that exploration of a new developmental pathway capable of rapid and parallel programming of a combinatorial library of nanoparticle-based delivery systems could lead to revolutionary breakthroughs in nanoparticle-based delivery systems.

Herein, we describe a rapid developmental pathway (Figure 1) that leverages the powers of (i) a combinatorial synthetic approach (Figure 1a) based on supramolecular assembly 17-19 and (ii) a digital microreactor<sup>20, 21</sup> (Figure 1b–d), towards the generation of a highly efficient nanoparticle-based gene delivery system. Unlike the slow, multistep syntheses employed for producing existing gene-delivery materials,<sup>3</sup> our supramolecular method (Figure 1a) enables a convenient, flexible, and modular method<sup>22</sup> for generating a combinatorial library of DNACSNPs, in which a broad structural/functional diversity covering the size variation, surface chemistry and DNA loading capacity was programmed into individual DNACSNPs by systematically altering the mixing ratios of five functional molecular building blocks (2-6), DNA plasmid (7a: enhanced green fluorescent protein (EGFP), and **7b**: firefly luciferase (FLuc)). In order to reduce human operational errors, accelerate handling procedures, enhance experimental fidelity, and achieve economical use of reagents, a digital Dual-Core Microreactor (DCM, Figure 1b-d) was designed and implemented to allow automated sampling, dilution, metering and mixing of 2–7, resulting in a combinatorial library composed of 648 different DNACSNPs within 2.5 h. The structural/functional diversity of the DNACSNPs library can be translated into diversity in performance by conducting transfection studies of individual DNACSNPs in 96 well plates containing mouse fibroblast cells. A small group of DNACSNPs that facilitates high levels of delivery performance was identified. We then carried out comprehensive characterizations on these DNACSNPs revealing that improved transfection performance can be attributed to the defined size, surface chemistry, zeta potential, uniformity and dynamic stability. Compared to the leading gene transfection reagents, such as lipofectamine 2000 and RGD-jet-PEI, the identified 40-nm TAT/RGD-DNACSNPs (1) with defined surface chemistry (RGD: $^{23-26}$  a  $\alpha_v \beta_3$  binding peptide and TAT $^{27-29}$  a cell-penetrating peptide) exhibited significantly improved gene transfection efficiency and low toxicity in a number of cancer cell lines and fibroblast cells.

# **Results and Discussion**

The five molecular building blocks (Figure 1a) – CD-PEI (2), Ad-PAMAM (3), Ad-PEG (4), RGD-PEG-Ad (5) and TAT-PEG-Ad (6) were prepared according to literatures, <sup>17, 22</sup> otherwise see Supporting Information. The functions of the five molecular building blocks are summarized below. First, the complementary CD-PEI (2) and Ad-PAMAM (3) are responsible for constructing cationic hydrogel networks that can encapsulate anionic DNA (7) forming the cores of DNACSNPs. Therefore, the DNA loading capacity of DNACSNPs is dependent on the net positive charges embedded in the hydrogel networks. By using electrophoresis analysis and ethidium bromide exclusion assay, <sup>22</sup> the nitrogen/phosphate (N/ P) ratio above five was determined. Second, Ad-PEG (4) serves as a capping/solvation reagent that not only constrains continuous growth of the DNA-encapsulated hydrogel networks, but also confers desired water solubility, structural stability, and passivation performance to the resulting DNACSNPs. Third, the two functional ligands (5 and 6) can be incorporated onto the surfaces of DNACSNPs via dynamic exchange in order to enable delivery specificity (to recognize a certain population of cells with  $\alpha_v \beta_3$ -integrin receptors) and cell transfusion capability (to foster internalization through membrane) of the resulting DNACSNPs, respectively. By systematically altering the mixing ratios among the five molecular building blocks (2-6) and DNA (7), distinct structural/functional properties (i.e., sizes and surface chemistry) can be programmed into individual DNACSNPs in the combinatorial library. It is important to point out that, in contrast to the lipid-like gene delivery system where the diversity was built upon the binary combination of a plethora of molecular precursors, <sup>30, 31</sup> the diversity of our DNACSNPs library was generated by ratiometric combination of only the five molecular building blocks and DNA loading.

It is feasible to manually prepare the DNACSNPs library by pipetting and mixing individual molecular building blocks and DNA plasmid at different ratios, while potential operation errors, slow handling speed, and a significant amount of sample consumption might compromise the throughput, fidelity and efficiency of the study. Digital microfluidic reactors <sup>32–35</sup> are promising platforms to overcome these challenges. In our case, an oil-free, digital DCM (Figure 1b–d) composed of (i) a Central Integrated Mixer (CIM) and (ii) an Auxiliary Integrated Mixer (AIM), was designed and implemented to systematically program the structural/functional diversity into the DNACSNPs libraries.

As illustrated in Figure 1b, there are eight parallel microchannels in CIM, capable of parallel generation of eight slugs with different mixing ratios of 2–7. Each microchannel was partitioned by hydraulic-actuated microvalves into nine confined plug regions, which were assigned for precise metering of five molecular building blocks (2–6), a DNA plasmid (7a: EGFP or 7b: FLuc) and phosphate buffered saline (PBS). AIM incorporates a pair of fractionally partitioned microchannels, capable of synchronized supply of functional ligands (5 and 6) at different concentrations. CIM and AIM were coupled together by a pair of degas modules (Supporting Information) that remove gas through liquid transferring processes.

By using the digital DCM, the automated preparation of DNACSNPs library was achieved by systematically modulating the mixing ratios of CD-PEI (2), RGD-PEG-Ad (5) and TAT-PEG-Ad (6) against fixed amounts of Ad-PAMAM (3), Ad-PEG (4) and plasmid DNA (7). In CIM, constant concentrations of 3 (3.0  $\mu$ M), 4 (60.0  $\mu$ M) and 7 (100 ng/ $\mu$ L) were assigned to fill into plug regions (ca. 20 nL per region). To adjust the quantities of CD-PEI (2), the eight uneven partitioned plug regions on the west side of CIM were utilized to accommodate different volumes of 2 (4.0  $\mu$ M, 10–80 nL). The concentration modulation of the two functional ligands (5 and 6) was accomplished in AIM, where each fractionally partitioned microchannel is responsible for ratiometric mixing of the respective ligand (5: 0–9.6  $\mu$ M and 6: 0–12.0  $\mu$ M) with PBS. In each operation cycle, CIM generate eight 200-nL

slugs containing different precursor mixtures, which were introduced into 96-well plates by nitrogen through eight poly(tetrafluoroethylene) (PTFE) tubes attached at the east side of CIM. In order to produce sufficient quantity of DNACSNPs for subsequent transfection studies, 50 cycles were performed continuously in CIM within 90 s, affording eight different DNACSNP solutions (ca. 10 μL each, 100 ng DNA). (See the supporting movie clip and Supporting Information). The resulting DNACSNPs libraries were incubated in the 96-well plates for 20 min at room temperature prior to transferring to the 96-well plates containing NIH 3T3 cells (ca. 8000 cells/well) for transfection studies in parallel. The gene transfection efficiency was evaluated by a plate reader after culturing the cells at 37°C (5% CO<sub>2</sub>) for 24 h.

In proof-of-concept trials a CMV promoter-driven EGFP-encoded plasmid DNA was encapsulated into the DNACSNPs. Before a full-scale screening with three variables (i.e., CD-PEI (2), RGD-PEG-Ad (5) and TAT-PEG-Ad (6)), simplified studies on two pairs of variables (i.e., CD-PEI (2) vs. RGD-PEG-Ad (5) and CD-PEI (2) vs. TAT-PEG-Ad (6)) were conducted in search of the optimal transfection performance of RGD-DNACSNPs and TAT-DNACSNPs, respectively. The transfection outcomes (96 data points in each case) were fitted into two 3-dimensional (3D) profiles (Figure 2a and b). The full-scale screening was accomplished by systematically programming the three variables, i.e., 8 different concentrations for CD-PEI (2), 9 for RGD-PEG-Ad (5) and 9 for TAT-PEG-Ad (6). Using the digital DCM, a combinatory library composed of 648 different DNACSNPs was generated within 2.5 h to fill up seven 96-well plates. To ensure the operation fidelity, all the experiments were conducted in triplicate. A 4D gene expression plot was employed to summarize the results from the full-scale screening of the combinatorial library revealing that the optimal transfection performance of DNACSNPs was achieved at CD-PEI (2) concentration of 0.6–0.8 µM, a RGD-PEG-Ad (5) of 0.24–0.48 µM and TAT-PEG-Ad (6) of 0.4–0.8 μM. We also note the transfection performance of the TAT/RGD-DNACSNPs (1a) is significantly improved compared with those observed for RGD-DNACSNPs and TAT-DNACSNPs. To validate the general applicability of this developmental pathway we performed a relatively smaller screening to identify DNACSNPs capable of highly efficient delivery of FLuc-encoded plasmid DNA. We manipulated three variables ( $5 \times 5 \times 5$ concentrations) resulting in a combinatorial library of 125 different TAT/RGD-DNACSNPs (1b). The consistent results (Figure 2d) were observed for the transfection performance of TAT/RGD-DNACSNPs (1b) for delivering FLuc-DNA plasmid. All four sets of screening studies suggested that the TAT/RGD-DNACSNPs (1) composed of CD-PEI (2,  $0.7 \pm 0.1$  $\mu$ M), RGD-PEG-Ad (5, 0.28  $\pm$  0.04  $\mu$ M) and TAT-PEG-Ad (6, 0.60  $\pm$  0.20  $\mu$ M) show the best transfection performance.

To understand how the identified synthetic variables contribute to the optimal transfection performance we scaled up the production of TAT/RGD-DNACSNPs (1a) under three sets of synthetic variables for characterization at three CD-PEI (2) concentrations (0.6, 0.8 and 1.2  $\mu$ M) while keeping concentrations of the RGD-PEG-Ad (0.28  $\mu$ M) and TAT-PEG-Ad (0.60  $\mu$ M) constant. We realize that CD-PEI (2) concentration affects the cross-linking degrees of the CD-PEI/Ad-PAMAM hydrogel cores in TAT/RGD-DNACSNPs (1), leading to the size variation. Both transmission electron microscopy (TEM, Figure 3a) and scanning electron microscopy (SEM, Figure 3b) were employed to examine the morphology and sizes of the resulting TAT/RGD-DNACSNPs (1). Three distinct sizes of  $42 \pm 4$ ,  $86 \pm 9$  and  $160 \pm 13$  nm were observed for those produced at CD-PEI (2) concentrations of 0.6, 0.8 and 1.2  $\mu$ M, respectively. Interestingly, we also noticed that the size distributions of the DCM-produced DNACSNPs are much narrower than those prepared manually (Tabe 1), and these results were confirmed by dynamic light scattering (Zetasizer Nano, Malvern Instruments Ltd) measurements (Figure 3c). We attribute the greater control upon DNACSNP size distribution to the precision and reproducibility of the sampling, metering and mixing

processes in the digital DCM. In addition, no significant size variations were observed by introducing different concentrations ranging from 0 to 1.2 µM of ligands (5 and/or 6). We also characterized the surface potentials of these TAT/RGD-DNACSNPs (Figure 3d), which spanned a range of 3.7±0.2–6.8±0.3 mV. In addition, the actual surface coverages of RGD-PEG-Ad (5) and TAT-PEG-Ad (6) were estimated by measuring absorption intensity of a FITC-labeled analog (i.e., FITC-PEG-Ad)<sup>22</sup> on the DNACSNPs prepared by the respective synthetic parameters (See Supporting Information). The results indicated that 0.28 and 0.60 μM of RGD-PEG-Ad (5) and TAT-PEG-Ad (6) reflect 5 and 9 % of surface coverage on the resulting DNACSNPs, respectively. In short, TAT/RGD-DNACSNPs (1) with sizes of 40 and 80 nm, as well as 5% RGD-PEG-Ad (5) and 9% TAT-PEG-Ad (6) coverages<sup>22</sup> exhibited optimal cell transfection performance. Finally, in order to understand the dynamic stability of 40- and 80-nm TAT/RGD-DNACSNPs (1) we employed real-time DLS measurements to monitor their size variation (i) at different pH values (pH 3.8-9.2) and temperatures (25, 37 and 60°C), (ii) in the presence of 10% serum, and (iii) at different storage times in presence of physiological salt concentrations (Figure 3e-h). The stability of the TAT/RGD-DNACSNPs (1) can be attributed to the multivalent CD/Ad recognition and electrostatic interactions between Ad-PAMAM/CD-PEI hydrogel and DNA. In order to study the role of Ad-PEG (4) for dynamic stability of DNACSNPs we monitored the size variation of DNACSNPs (Figure 3i and j) by removing the excess amount of Ad-PEG (4) in the mixture through membrane dialysis (MWCO 10kD) under pH 5.5 and 7.2. It was found that the 40- and 80-nm TAT/RGD-DNACSNPs (1) were disassembled gradually within 6 h at both pH conditions. In addition, the TAT/RGD-DNACSNPs (1) exhibited faster disassembly behavior at pH 5.5 (< 2 h) than at pH 7.2, which allows the TAT/RGD-DNACSNPs (1) efficiently degraded inside the endosomes and released the DNA intracelluarly.<sup>36</sup>

To compare the transfection performance of the optimal 40-nm TAT/RGD-DNACSNPs (1) with the leading transfection reagents (i.e., RGD-jet-PEI and lipofectamine 2000) we carried out the dose-dependent gene transfection studies in 24-well plates with encapsulated-DNA (50 and 1000 ng per well) by using a collection of cells, including NIH 3T3 (Mouse fibroblast cell line), HeLa (human cervix epithelial carcinoma cell line), A549 (human lung cancer cell line), U87 (human glioblastoma cell line), MCF7 (human breast adenocarcinoma cell line), PC3 (human prostate cancer cell line), and IMR-90 (human fibroblast cell line). The gene transfection studies results (Figure 4a and b) indicated that the 40-nm TAT/RGD-DNACSNPs (1) exhibited significantly improved transfection performance compared to those observed for RGD-jet-PEI and lipofectamine 2000 across the different cancer and fibroblast cell lines at high dosage of DNA (1000 ng DNA per well). Higher than 70% transfection efficiencies were observed across various cancer cell lines we tested, even for the PC3 cell line that is difficult to be transfected by those two commercial reagents. IMR-90, one of the hard-to-transfect human fibroblast cell lines, was tested and the transfection efficiency of DNACSNPs toward IMR-90 cells reached to 17±7 %, which can be visualized by the fluorescence micrographs (Figure 4c). The transfection efficiency of 40nm TAT/RGD-DNACSNPs (1) for IMR-90 cells is 11-fold and 5-fold higher than that of RGD-jet-PEI and lipofectamine 2000, respectively. It is important to note that 40-nm TAT/ RGD-DNACSNPs (1) still achieved high transfection efficiency at low dosage of DNA (50 ng per well), indicating the outstanding gene delivery ability of DNACSNPs. Moreover, the cell viability assay results indicated that the 40-nm TAT/RGD-DNACSNPs (1) exhibited negligible cytotoxicity. More than 96 ± 3 % of DNACSNPs-transfected cells showed normal viability even in the presence of high dosage of reagent (1000 ng DNA per well). In contrast, at high DNA dose more than 34% and 22% NIH 3T3 cells died after treatment by RGD-jet-PEI and lipofectamine 2000 for 48 h (Figure 4d).

# **Experimental Section**

#### General

Reagents and solvents were purchased from Sigma-Aldrich (St. Louis, MO) and used as received without further purification or otherwise noted. Branched polyethylenimine (PEI, MW = 10 kD) was purchased from Polysciences Inc (Washington, PA). Polymers contain primary, secondary and tertiary amine groups in approximately 25/50/25 ratio. 1stgeneration polyamidoamine dendrimer (PAMAM) with 1, 4-diaminobutane core and amine terminals in 20% wt methanol solution was purchased from Dendritic Nanotechnologies, Inc (Mount pleasant, MI). 1-Adamantanamine (Ad) hydrochloride and β-cyclodextrin (β-CD) were purchased from TCI America (San Francisco, CA). N-hydroxysuccinimide (SCM) and maleimido (MAL) hetero-functionalized polyethylene glycol (SCM-PEG-MAL, MW = 5 kD) was obtained from NANOCS Inc (New York, NY). Phosphate-buffered saline (PBS, 1X, pH 7.2  $\pm$  0.05) for sample preparation. 6-Mono-tosyl-β-cyclodextrin (6-OTs-β-CD) was prepared following the literature reported method.<sup>37</sup> Octa-Ad-grafted polyamidoamine dendrimer (Ad-PAMAM), CD-grafted branched polyethylenimine (CD-PEI) and Ad-grafted polyethylene glycol (Ad-PEG) were prepared as the method we reported before. <sup>17</sup> Dry CH<sub>2</sub>Cl<sub>2</sub> was obtained by refluxing over CaH<sub>2</sub> and freshly distilled before use. NIH 3T3 (mouse embryonic fibroblast cell line), HeLa (human cervix epithelial carcinoma cell line), A549 (human lung cancer cell line), U87 (human glioblastoma cell line), MCF7 (human breast adenocarcinoma cell line), PC3 (human prostate cancer cell line) and IMR-90 (human primary fibroblasts cell line) were purchased from American Type Culture Collection (ATCC). The Dulbecco's Modified Eagle Medium (DMEM), Earl's Modified Eagle's Medium (EMEM) growth medium, RPMI-1640, Opti-MEM reduced serum medium and Penicillin/streptomycin were obtained from Invitrogen (Carlsbad, CA). Fetal Bovine Serum (FBS) and EGFP-encoded plasmid DNA (pMAX EGFP®, 4.3 kb) and luciferase-encoded plasmid DNA (FLuc-DNA, 6.6 kb) were obtained from Lonza Walkerrsville Inc (Walkerrsville, MD). 4', 6-Diamidino-2-phenylindole (DAPI) was purchased from Invitrogen (Carlsbad, CA). RGD (RGDC) and TAT peptides (CGRKKRRORRR) were purchased from GenScript Corp. (Piscataway, NJ).

<sup>1</sup>H NMR spectra were recorded on a Bruker Avance 400 spectrometer in deuterated solvents. Mass spectra were acquired using an Applied Biosystems Voyager DE-STR MALDI-TOF mass spectrometer (Framingham, MA). Dynamic light scattering and zeta potentials of DNA⊂SNPs (1) were measured on Zetasizer Nano instrument (Malvern Instruments Ltd., United Kingdom). Transmission electron microscope (TEM) images were measured on Philips CM 120 electron microscope operating with an acceleration voltage of 120 kV. SEM images of DNA⊂SNPs (1) were obtained with a JEOL JSM-6700F SEM. Freeze-dried samples on a silicon surface were sputter-coated with gold before measurement. Cell imaging and gene transfection studies were performed on a Nikon TE2000S inverted fluorescent microscope with a CCD camera (Photomatrix, Cascade II), X-Cite 120 Mercury lamp, automatic stage, and filters for three fluorescent channels (W1 (DAPI), W2 (EGFP and AO) and W3 (PI)).

#### The fabrication and operations of Digital Dural Core Microreactor (DCM)

The fabrication and operations of Digital Dural Core Microreactor (DCM) are described in the Supporting Information.

# Programming supramolecular nanoparticles in a digital dual-core microreactor

By using the digital DCM, two 3D and two 4D screenings were performed sequentially. In Figure 2a, 96 different RGD-DNACSNPs were generated by altering concentrations of CD-PEI  $(0.2, 0.4, 0.6, 0.8, 1.0, 1.2, 1.4 \text{ and } 1.6 \mu\text{M})$  and RGD-Ad-PEG (0, 0.08, 0.16, 0.24,

0.32, 0.40, 0.48, 0.56, 0.64, 0.72, 0.80, 0.88 and 0.96 μM). In Figure 2b, 96 different TAT-DNACSNPs were generated by altering concentrations of CD-PEI (0.2, 0.4, 0.6, 0.8, 1.0, 1.2, 1.4 and 1.6 μM) and TAT-Ad-PEG (0, 0.10, 0.20, 0.30, 0.40, 0.50, 0.60, 0.70, 0.80, 0.90, 1.00, 1.10 and 1.20 μM). In Figure 2c, 648 different TAT/RGD-DNACSNPs (**1a**) were generated by altering concentrations of CD-PEI (0.2, 0.4, 0.6, 0.8, 1.0, 1.2, 1.4 and 1.6 μM), TAT-Ad-PEG (0, 0.10, 0.20, 0.30, 0.40, 0.60, 0.80, 1.00, and 1.20 μM) and RGD-Ad-PEG (0, 0.08, 0.16, 0.24, 0.32, 0.48, 0.64, 0.80, and 0.96 μM). In Figure 2d, 125 different TAT/RGD-DNACSNPs (**1b**) were generated by altering concentrations of CD-PEI (0.4, 0.6, 0.8, 1.0 and 1.2 μM), TAT-Ad-PEG (0, 0.10, 0.30, 0.60 and 1.20 μM) and RGD-Ad-PEG (0, 0.16, 0.24, 0.32 and 0.48 μM).

## **Dynamic light scattering (DLS)**

DLS experiments were performed with a Zetasizer Nano instrument (Malvern Instruments Ltd., United Kingdom) equipped with a 10-mW helium-neon laser ( $\lambda = 632.8$  nm) and thermoelectric temperature controller. Measurements were taken at a 90° scattering angle. The sizes and the standard derivations of assembled TAT/RGD-DNACSNPs (1) were calculated by averaging the values of at least three measurements.

#### Transmission electron microscope (TEM)

The morphology and sizes of TAT/RGD-DNA $\subset$ SNPs were directly examined using transmission electron microscope. The studies were carried out on a Philips CM 120 electron microscope, operating at an acceleration voltage of 120 kV. The TEM samples were prepared by drop-coating 2  $\mu$ L of TAT/RGD-DNA $\subset$ SNPs solutions onto carbon-coated copper grids. Excess amounts of slugs were removed by filter papers after 45 s. Subsequently, the surface-deposited TAT/RGD-DNA $\subset$ SNPs were negatively stained with 2% uranyl acetate for 45 s before the TEM studies.

# Scanning electron microscope (SEM)

We prepared the TAT/RGD-DNA $\subset$ SNPs samples for SEM observation by a standard procedure: Briefly, 2  $\mu$ L of TAT/RGD-DNA $\subset$ SNPs solution was drop-coated onto silica substrate and freeze-dry by dry ice. Then, the samples were sputter coated with gold before examination with a Hitachi S800 field emission SEM at an accelerating voltage of 10 keV.

## Zeta potential measurements

Zeta potentials of TAT/RGD-DNACSNPs (1) were determined by photon correlation spectroscopy using a Zetasizer Nano instrument, (Malvern Instruments, Malvern, Worcestershire, UK). The measurements were performed at 25°C with a detection angle of 90°, and the raw data were subsequently correlated to Z average mean size using a cumulative analysis by the Zetasizer software package.

# Stability studies of 40 and 80-nm TAT/RGD-DNA⊂SNPs

# Stability of 40 and 80-nm TAT/RGD-DNACSNPs in different pH values—

Information on the stability of TAT/RGD-DNACSNPs in variable pH environments is needed to understand its characteristics under physiological conditions and to provide a reasonable pH range for chemical modification of TAT/RGD-DNACSNPs for further experiments. The pH-dependent stability of TAT/RGD-DNACSNPs as a function of pH was tested at a range from 3.8 to 9.2 by DLS analysis. The stock solutions of 40- and 80-nm TAT/RGD-DNACSNPs were prepared in PBS buffer containing monobasic potassium phosphate (I = 1.5 mM), sodium chloride (I = 155 mM), and dibasic sodium phosphate (I = 2.7 mM). The following buffers were used to adjust the pH values of the TAT/RGD-DNACSNPs samples in the stock solution: 100 mM HCl-KCl buffer (pH 0–2.0), 100 mM

Glycine-HCl buffer (pH 2.2–3.6), 100 mM CH<sub>3</sub>COOH-CH<sub>3</sub>COONa buffer (pH 3.7–5.6), 100 mM Na<sub>2</sub>HPO<sub>4</sub>-NaH<sub>2</sub>PO<sub>4</sub> buffer (pH 5.8–8.0), 100 mM Tris-HCl buffer (pH 7.0–9.0) and 100 mM Na<sub>2</sub>CO<sub>3</sub>-NaHCO<sub>3</sub> buffer (pH 9.2–10.8). Typically, a 100- $\mu$ L portion of TAT/RGD-DNACSNPs stock solution was mixed with a 900- $\mu$ L portion of different pH value buffer solutions (I = 100 mM). The resulting solutions were shaken and equilibrated until the pH stabilized. The final pH values were determined by pH meter.

# Stability of 40 and 80-nm TAT/RGD-DNACSNPs under different temperatures —To understand the thermal stability of the TAT/RGD-DNACSNPs we employed real-time DLS measurements to monitor the size varieties of both of the 40, and 80 nm TAT/RGD.

DLS measurements to monitor the size variation of both of the 40- and 80-nm TAT/RGD-DNACSNPs in PBS at 25, 37 and 60 °C. In each case the samples were equilibrated under a specific temperature for 20 min prior to data acquisitions.

**Stability of 40 and 80-nm TAT/RGD-DNA**CSNPs in 10% Serum—To understand the stability of the TAT/RGD-DNACSNPs in presence of serum we employed DLS measurements to observe the size variation of both of the 40- and 80-nm TAT/RGD-DNACSNPs in the mixture of serum and PBS (1:9, V/V). The samples were kept at room temperature for 4h prior to data acquisitions.

Stability of 40 and 80-nm TAT/RGD-DNA⊂SNPs under a physiological ionic strength—To ensure the *in vivo* stability of the TAT/RGD-DNA⊂SNPs it is critical to examine the size variation of them under a physiological ionic strength. The 40- and 80-nm TAT/RGD-DNA⊂SNPs were prepared in PBS solutions (pH = 7.2). We employed real-time DLS measurements to monitor the size variation of both of the 40- and 80-nm TAT/RGD-DNA⊂SNPs at different times. The TAT/RGD-DNA⊂SNPs sizes were recorded for 48 h.

#### Cell culture

NIH 3T3, HeLa, A549, U87 and IMR-90 cell lines were routinely maintained in DMEM containing 10% fetal bovine serum (FBS) and 1% penicillin/streptomycin (Invitrogen, Carlsbad, CA). MCF7 was cultured in EMEM containing 10% fetal bovine serum (FBS) and 1% penicillin/streptomycin. PC3 was cultured in RPMI-1640 containing 10% fetal bovine serum (FBS) and 1% penicillin/streptomycin.

#### Gene transfection studies

Cells ( $5\times10^4$  cells/well) were plated in 24-well plates and allowed to adhere overnight. EGFP-encoded DNA was diluted in  $1\times$  TE buffer. The 40-nm TAT/RGD-DNACSNPs were prepared on DCM and incubated for 20 min at room temperature before transfection experiments. The 40-nm TAT/RGD-DNACSNPs in PBS ( $10~\mu L$ ) was diluted with  $100~\mu L$  Opti-MEM medium and subsequently transferred to each well. For the control groups, RGD-jet-PEI and lipofectamine 2000 were used as a standard transfection reagent and operated according to the protocol provided by the manufacturers. TAT/RGD-DNACSNPs along with controls were incubated with the cells for 4 hours then removed by aspirating, and replaced with  $500~\mu L$ /well of fresh culture media. Cells were allowed to grow for 24 h at  $37^{\circ}C$  and 5% CO<sub>2</sub> and then fixed (4% paraformaldehyde for 15 min at room temperature), then washed with PBS three times, stained with DAPI, and finally rinsed with PBS prior to EGFP expression analysis by fluorescence microscope.

## Conclusion

In conclusion, we have demonstrated a rapid developmental pathway toward generation of a highly efficient gene delivery system by leveraging the powers of a supramolecular synthetic approach and a custom-designed digital microreactor. This pathway can be

adopted for the development of nanoparticle-based vectors capable of delivering a variety of loads, such as gene, drugs, proteins and their mixtures. We are currently exploring the use of the DNACSNP-based transfection reagents for reprogramming of human primary fibroblast cells in order to generate induced pluripotent stem cells that is crucial in the field of regulative medicine.<sup>38</sup>

# Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

# **Acknowledgments**

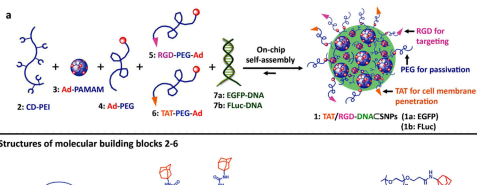
This research was supported by NIH-NCI NanoSystems Biology Cancer Center (U54CA119347), R21 grant (EB008419–01) and California Institute of Regenerative Medicine (RT1-01022-1).

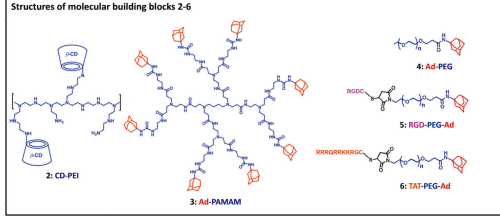
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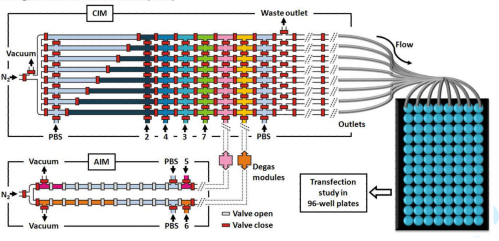
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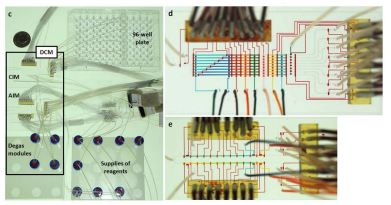
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## b Digital Dual-Core Microreactor (DCM)





#### Figure 1.

A rapid developmental pathway that leverages the powers of (i) a combinatorial synthetic approach based on supramolecular assembly and (ii) an automated microreactor has been demonstrated for screening and production of a highly efficient nanoparticle-based gene delivery system. a) Graphical schematic representations of the self-assembly approach for producing a combinatorial library of DNA encapsulated supramolecular nanoparticles (DNACSNPs), in which a broad structural/functional diversity can be programmed into individual DNACSNPs (1) by systematically altering the mixing ratios of the five functional molecular building blocks i.e., CD-PEI (2), Ad-PAMAM(3), Ad-PEG (4), RGD-PEG-Ad (5) and TAT-PEG-Ad (6), as well as DNA plasmid (7a: enhanced green fluorescent protein (EGFP) and 7b: firefly luciferase (FLuc)). b) Graphical illustration of a digital Dual-Core Microreactor (DCM). The device settings are composed of a central integrated mixer (CIM) and an Auxiliary Integrated Mixer (AIM), and degas modules bridges. The operation of the circuit was computer controlled using color-coded pressure-driven valves: red - positive pressure, off/on; yellow – peristaltic pumping; green – vacuum. Eight slugs containing eight different formulated DNA encapsulated DNACSNPs were generated in parallel and transferred to a 96-well plate through PTFE tubes. The gene transfection performance for each combination was evaluated by a plate reader. c) The picture of the whole set of actual instrument. d and e) Optical images of the CIM and AIM, respectively. The various channels were loaded with dyes to visualize the different components.

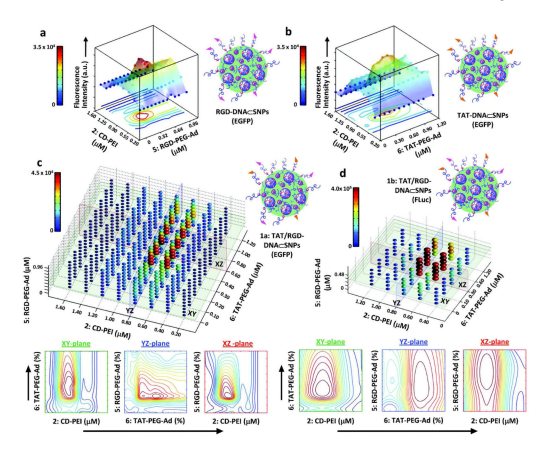


Figure 2.
a) A 3D profile of gene transfection performance of RGD-DNACSNPs (EGFP) with variation of CD-PEI (2) and RGD coverage (96 data points). b) A 3D profile of gene transfection performance of TAT-DNACSNPs (EGFP) with variation of CD-PEI (2) and TAT coverage (96 data points). c) A 4D profile of gene transfection performance of TAT/RGD-DNACSNPs (EGFP) with variation of CD-PEI (2), TAT and RGD coverages (648 data points). The XY, YZ and XZ plates across the best performance were simplified and denoted 2D contour images. d) A 4D profile of gene transfection performance of TAT/RGD-DNACSNPs (FLuc) with variation of CD-PEI (2), TAT and RGD coverages (125 data points). The XY, YZ and XZ plates across the best performance were simplified and denoted 2D contour images.

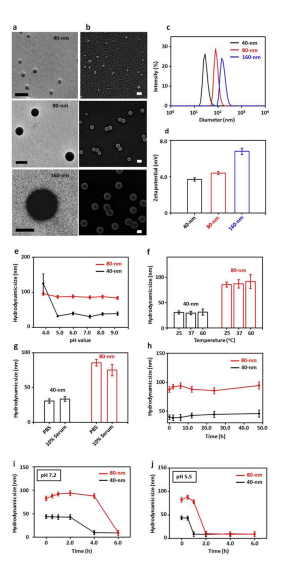


Figure 3. a) and b) TEM and SEM images of the resulting TAT/RGD-DNA $\subset$ SNPs with different sizes of  $42 \pm 4$ ,  $86 \pm 9$  and  $160 \pm 13$  nm. Scale bars: 100 nm. c) Dynamic light scattering (DLS) was employed to measure DNA $\subset$ SNPs hydrodynamic sizes in PBS buffer. d) Zeta potentials variations of DNA $\subset$ SNPs in PBS buffer. Stability studies of 40- and 80-nm DNA $\subset$ SNPs under different conditions: e) pH-dependent size variations of DNA $\subset$ SNPs in the respective buffer solutions with pH values ranging from 3.8 to 9.2. Error bars are obtained from three measurements. f) Temperature-dependent size variation of DNA $\subset$ SNPs in PBS buffer (pH = 7.2) with different temperatures at 25, 37 and 60 °C. g) Size variations of DNA $\subset$ SNPs in presence and absence of 10% serum containing PBS buffer after 4 h incubation. Error bars are obtained from three measurements. h) Time-dependent size variation of DNA $\subset$ SNPs from 0–48 h in PBS (pH 7.2). i and j) Time-dependent dynamic stability of DNA $\subset$ SNPs under dialysis at pH 5.5 and 7.2, respectively.

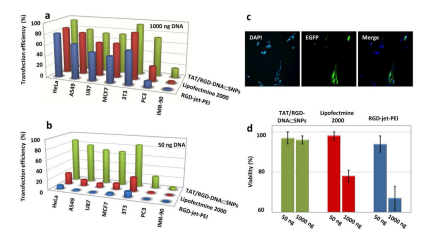


Figure 4.
a) and b) Transfection efficiencies of the optimal 40-nm TAT/RGD-DNACSNPs at high (1000 ng DNA per well) and low (50 ng DNA per well) DNA dosage along with control delivery reagents (lipofectamine 2000 and RGD-jet-PEI) for a collection of cell lines, including NIH 3T3 (Mouse fibroblast cell line), HeLa (human cervix epithelial carcinoma cell line), A549 (human lung cancer cell line), U87 (human glioblastoma cell line), MCF7 (human breast adenocarcinoma cell line), PC3 (human prostate cancer cell line) and IMR-90 (human fibroblast cell line) in 24-well plates. c) The representative fluorescence micrographs of IMR-90 cells transfected by 40-nm TAT/RGD-DNACSNPs at high dosage of DNA (1000 ng DNA/well). d) Cytotoxicity of RGD-jet-PEI, lipofectamine 2000 and 40-nm TAT/RGD-DNACSNPs at high and low dosage of DNA transfected NIH 3T3 cells determined by cell viability assay after 48 h of transfection.

Table 1

Comparison of TAT/RGD-DNA⊂SNPs synthesized by DCM and conventional pipetting

	DCM	Pipetting <sup>22</sup>
Throughput	250 conditions/h	<20 conditions/h
Size distribution	Excellent (PDI <sup>a</sup> : <0.05)	Good (PDI <sup>a</sup> : 0.10~0.20)
Operation error	No	Possible
Reproducibility	High	Modest

 $<sup>^{\</sup>it a}{\rm PDI}{:}$  The polydispersity index obtained from DLS measurements.