Physicochemical and biological characterization of targeted, nucleic acid-containing nanoparticles

Derek W. Bartlett and Mark E. Davis
Chemical Engineering, California Institute of Technology, Pasadena, California 91125, USA

Abstract

Nucleic acid-based therapeutics have the potential to provide potent and highly specific treatments for a variety of human ailments. However, systemic delivery continues to be a significant hurdle to success. Multifunctional nanoparticles are being investigated as systemic, nonviral delivery systems, and here we describe the physicochemical and biological characterization of cyclodextrin-containing polycations (CDP) and their nanoparticles formed with nucleic acids including plasmid DNA (pDNA) and small interfering RNA (siRNA). These polycation/nucleic acid complexes can be tuned by formulation conditions to yield particles with sizes ranging from 60–150 nm, zeta potentials from 10–30 mV, and molecular weights from ~7×10^7–1×10^9 g mol^-1 as determined by light scattering techniques. Inclusion complexes formed between adamantane (AD)-containing molecules and the β-cyclodextrin molecules enable the modular attachment of polyethylene glycol (AD-PEG) conjugates for steric stabilization and targeting ligands (AD-PEG-transferrin) for cell-specific targeting. A 70-nm particle can contain ~10,000 CDP polymer chains, ~2,000 siRNA molecules, ~4,000 AD-PEG_{5000} molecules, and ~100 AD-PEG_{5000}-Tf molecules; this represents a significant payload of siRNA and a large ratio of siRNA to targeting ligand (20:1). The particles protect the nucleic acid payload from nuclease degradation, do not aggregate at physiological salt concentrations, and cause minimal erythrocyte aggregation and complement fixation at the concentrations typically used for in vivo application. Uptake of the nucleic acid-containing particles by HeLa cells is measured by flow cytometry and visualized by confocal microscopy. Competitive uptake experiments show that the transferrin-targeted particles display enhanced affinity for the transferrin receptor through avidity effects (multi-ligand binding). Functional efficacy of the delivered pDNA and siRNA is demonstrated through luciferase reporter protein expression and knockdown, respectively. The analysis of the CDP delivery vehicle provides insights that can be applied to the design of targeted nucleic acid delivery vehicles in general.

Keywords

Nanoparticles; transferrin targeting; plasmid DNA; siRNA

INTRODUCTION

Nucleic acid-based therapeutics are envisioned to play a significant role in the next generation of treatments for a variety of diseases such as cancer. In addition to the classic gene therapy approach of delivering DNA to replace mutated or absent genes, nucleic acid molecules can also be used to regulate the production of disease-associated proteins at both the transcriptional and translational levels. These nucleic acid-based drugs have received significant attention as
promising new therapeutics, yet their application \textit{in vivo} has been largely limited by the challenge of delivery; this has been particularly true for systemic delivery.

Naked nucleic acid molecules are rapidly degraded by ubiquitous nucleases present in the bloodstream. Double-stranded nucleic acid molecules ranging in size from small-interfering RNAs (siRNAs) to plasmids (pDNA) have a half-life of less than one hour in serum (1–3). Selective chemical modification of nucleic acids can increase nuclease resistance and enable systemic delivery of naked siRNA molecules with functional efficacy \textit{in vivo} (1,4). However, even nuclease-stabilized nucleic acids must still overcome other elimination barriers such as renal clearance that severely limit the efficacy of systemically delivered, small nucleic acid therapeutics (5). Attachment of specific targeting ligands can induce binding to protein carriers or uptake by the desired population of cells to be treated. Bioconjugates of the nucleic acid therapeutics covalently attached directly to targeting ligands such as cholesterol and antibodies have shown efficacy both \textit{in vitro} and \textit{in vivo} (4,6). While these methods for nuclease stabilization and covalent attachment of targeting ligands are promising for small nucleic acid therapeutics, the use of lipid- or polymer-based delivery vehicles is an approach for systemic delivery that can provide functions not achievable with naked nucleic acids or their covalent attachment to targeting moieties.

Carrier-mediated delivery has several advantages over the delivery of individual nucleic acid molecules. Encapsulation of the payload within a lipid bilayer or through electrostatic interactions is nonspecific, so these delivery vehicles can be used for generalized nucleic acid delivery. The use of a carrier enables delivery of many nucleic acid molecules per uptake event (this is especially important if the uptake involves highly specific cell-surface receptors since they are typically low in number), and isolation from exposure to the systemic environment can permit the use of unmodified nucleic acids (7). Modularly designed delivery vehicles can also take advantage of covalent or non-covalent attachment of hydrophilic polymers for steric stabilization and/or targeting ligands for cell-specific delivery, two critical features for systemic delivery (7,8). Such modifications can affect the resulting biodistribution of delivery vehicles through passive and/or active targeting (7–10).

Passive targeting occurs as a result of the intrinsic physicochemical properties of the delivery vehicle. For example, the charge and size of the delivery vehicle alone can bias its biodistribution. The charge of the delivery vehicle significantly impacts its interaction with components of the bloodstream; highly charged particles can lead to complement activation, while near-neutral particles exhibit reduced phagocytic uptake (11,12). Specifically, cationic polymers such as polylysine and polyethylenimine have been shown to activate the complement system, and increasing polycation length and surface charge density lead to higher complement activation (11). Rapid binding of charged molecules by complement proteins or other opsonins can lead to immune stimulation and rapid clearance of the delivery vehicles from the bloodstream. The size of the delivery vehicle also matters for systemic delivery. Based on measured sieving coefficients for the glomerular capillary wall, it is estimated that the threshold for first-pass elimination by the kidneys is approximately 10 nm (diameter) (13), placing a lower size limit on the assembled delivery vehicles. On the other end of the size spectrum, macromolecular complexes preferentially accumulate in tumors through the enhanced permeability and retention (EPR) effect. However, large macromolecules or delivery vehicles exhibit limited diffusion in the extracellular space, such as the tumor interstitium, and in the complex intracellular environment (14); in both situations, restricted movement will severely limit efficacy by preventing uptake by a sufficient number of cells or hindering the ability of the delivered particles to localize to intracellular compartments such as the nucleus. Other restrictions limiting the maximum size of delivery vehicles can also be imposed by the selectivity of receptors on certain cell types. For example, a study by Rensen \textit{et al.} demonstrated that particles larger than 70 nm in diameter were not recognized by the asialoglycoprotein
Recent efforts toward targeted delivery have focused on decorating the surface of delivery vehicles with cell surface receptor-specific targeting ligands as a means of active targeting. Hydrophilic polymers, such as polyethylene glycol (PEG), can be attached to the surface of the delivery vehicles to mask surface charge and prevent nonspecific interactions, helping to prevent unwanted binding to components of the bloodstream, slow uptake by the reticuloendothelial system (RES), and alter the cellular uptake patterns (8,16). Further addition of entities that can interact with cell surface receptors such as the receptors’ natural ligands, antibodies, or antibody fragments allows the delivery vehicle to target particular cell types and undergo receptor-mediated endocytosis upon binding to the surface receptor (7,8,15).

In light of these considerations, a successful delivery vehicle must be engineered to have the following characteristics: (i) be small enough to extravasate and exhibit adequate tissue penetration, yet avoid rapid renal clearance; (ii) minimize nonspecific interactions and opsonization while providing specific targeting to a given cell; and (iii) protect the nucleic acid from degradation, but willingly release it upon arrival at the proper site. Over the past few years, we have been developing a synthetic delivery system based on a cyclodextrin-containing polycation (CDP) that has demonstrated some success in delivering nucleic acid payloads that include pDNA, siRNA, and DNAzymes (7,17–20). This delivery system is the first to be de novo designed for systemic delivery of nucleic acids and completely formulated by self-assembly (17). Here, we describe the physicochemical and biological characterization of the cyclodextrin-containing polycation delivery system and its formulation with nucleic acids. We illustrate issues of importance when designing any polycation composite with nucleic acids through the use of the CDP and point out when the conclusions are specific to this system only.

EXPERIMENTAL PROCEDURES

Formulation of nucleic acid particles

The chemical structure of the cyclodextrin-containing polycation is shown in Figure 1A. This short, linear polycation can be synthesized with (CDP-Im) or without (CDP) the imidazole groups on the terminal amines (17, 18). A schematic showing particle formation using CDP-Im and nucleic acid is shown in Figure 1B; particles are formed by mixing equal volumes of CDP-Im and nucleic acid. The ratio of positive (+) charges (2 moles of positive charge per CDP-Im monomer; denoted β-CD) to negative (−) charges (1 mole of negative charge per nucleotide) is defined as the formulation charge ratio (+/−). Polyethylene glycol (PEG) molecules containing adamantane (AD) on the proximal end and either methoxy (AD-PEG) or a targeting ligand such as transferrin (AD-PEG-Tf) on the distal end can be attached to the surface of the particles via inclusion complex formation between adamantane and the β-CD molecules on the polycation backbone (16, 17). The molecular weight of the PEG chain is typically 5,000 daltons (PEG5000).

Formulation of PEGylated/targeted nucleic acid particles

Pre-complexation (self-assembly)—Before addition to the nucleic acid, the CDP or CDP-Im was mixed with an AD-PEG conjugate at a 1:1 AD-PEG:β-CD (mol:mol) ratio in water. Targeted particles also require the addition of ligand-modified AD-PEG-X (e.g., AD-PEG-Tf (7,19,20)) as a percentage of the total AD-PEG in the mixture. For example, 1 mol% AD-PEG-Tf particles contain 0.01 moles AD-PEG-Tf for every 0.99 moles AD-PEG. The mixture of CDP (or CDP-Im), AD-PEG, and AD-PEG-Ligand in water was then added to an equal volume of nucleic acid in water such that the ratio of positive charges from CDP or CDP-Im to negative charges from the nucleic acid was equal to the desired charge ratio. A schematic
of this assembly process is shown in Figure 1B. Unless specified otherwise, all PEGylated or targeted nucleic acid particles used in this manuscript were prepared through the pre-complexation method.

**Post-complexation**—Particles were initially formulated in water by mixing equal volumes of nucleic acid and the cyclodextrin-containing polycation. After particles had formed, the AD-PEG and AD-PEG-Tf conjugates were added directly to the formulation mixture at the desired ratio of AD-PEG:β-CD (mol:mol). A schematic of this assembly process is shown in Figure 1B.

**Electrophoretic mobility shift assay**

siRNA-containing particles were formulated at different charge ratios by changing the amount of CDP added to 1 μg of siRNA. CDP was first dissolved in 10 μL water and then added to an equal volume of water containing 1 μg of nucleic acid. After a 30-min incubation at room temperature, 10 μL of each formulation was run on a 1% agarose gel and visualized by ethidium bromide staining.

**Individual particle charge ratio**

siRNA particles were formulated in 40 μL water at charge ratios from 5 to 30 (+/−). After formulation, the particles were separated from the free components by addition of 400 μL PBS, to cause particle aggregation, followed by centrifugation to pellet the aggregated particles. Since the CDP is terminated by primary amines, quantitation of polycation content was accomplished by measuring the amount of primary amines as follows. 400 μL of the supernatant was removed and combined with 200 μL of 0.01% w/v 2,4,6-trinitrobenzenesulfonic acid (Sigma). After incubating these samples for 2 h at 37°C, 200 μL 10% SDS and 100 μL of 1 N HCl were added to each sample before measuring the absorbance at 335 nm with a spectrophotometer. The amount of CDP in each sample was determined by comparison to a standard curve of CDP. The total positive charge present in each sample was calculated from the mass of CDP present using the fixed charge density of 0.0014 moles “+” per gram. This gave the number of unbound “+” charges present, so the number of bound “+” charges in the particles could be determined by subtracting the number unbound from the total “+” charges added during formulation. Assuming 100% incorporation of the nucleic acid into the particles at a formulation charge ratio of 3 (+/−) (based on the electrophoretic mobility shift assays; see Results), the ratio of CDP to nucleic acid within each particle is equal to the number of bound “+” charges divided by the total number of “−” charges from the nucleic acid.

**Serum stability**

Particles were formulated in water at a charge ratio of 3 (+/−) with an siRNA concentration of 0.05 g L−1. 10 μL of 100% mouse serum (Sigma) were added to 10 μL of the particle formulation and subsequently incubated for 4 h at 37°C and 5% CO2. 0.25 μg naked siRNA in 5 μL water were added to 5 μL of 100% mouse serum and also incubated for 4 h at 37°C and 5% CO2. For comparison to the t = 4 h samples, identical amounts of naked siRNA or siRNA particles were exposed to 50% mouse serum immediately before gel loading (t = 0 h). 10 μL of each sample (containing 0.25 μg siRNA) were loaded per well of a 1% agarose gel. Displacement of the nucleic acid from the particles was achieved by adding 1% sodium dodecyl sulfate (SDS) to the sample immediately prior to gel loading. Gel electrophoresis was performed by applying 100 V for 30 min, and the bands were visualized by ethidium bromide staining.
Dynamic light scattering (DLS)

Particle formulations were diluted to a volume of 1.4 mL, placed in a cuvette, and inserted into a ZetaPALS (Brookhaven Instruments Corporation) instrument to measure both the size and zeta potential. Reported effective hydrodynamic diameters represent the average values from a total of 5–10 runs of 30 seconds each, while zeta potentials represent the average of 10 runs each.

Transmission electron microscopy (TEM)

Particles containing CDP-Im and siRNA (0.1 g L$^{-1}$) and PEGylated particles containing CDP-Im, AD-PEG (1:1 AD-PEG:β-CD mole ratio), and siRNA (0.5 g L$^{-1}$) were formulated in water at a charge ratio of 3 (+/−). Samples were stained with 2% uranyl acetate and then examined with an EM201C electron microscope (Philips).

Atomic force microscopy (AFM)

Particles containing CDP-Im and siRNA (0.1 g L$^{-1}$) and PEGylated particles containing CDP-Im, AD-PEG (1:1 AD-PEG:β-CD mole ratio), and siRNA (0.5 g L$^{-1}$) were formulated in water at a charge ratio of 3 (+/−). 20 μL of each formulation solution were dropped on a freshly cleaved mica disc (Ted Pella, Inc.) and dried with pressurized air. Images were acquired with a Digital Instruments MultiMode AFM with a Nanoscope IV controller in tapping mode at a scan rate of 1 Hz using a BS Multi75 probe (BudgetSensors) with a resonant frequency of 75 kHz and a force constant of 3 N m$^{-1}$. Height images were flattened and processed for visualization with the derivative matrix convolution filter using WSxM scanning probe microscopy software (Nanotec Electronica).

Isothermal titration calorimetry (ITC)

A MicroCal MCS titration calorimeter was used to investigate the thermodynamic properties of the interaction between AD-PEG conjugates and the β-cyclodextrin molecules on the CDP-Im backbone. CDP-Im (free or in particles) at 0.22 mM total β-CD in water was placed in the sample cell of the instrument. The reference cell contained water alone without CDP-Im. Small amounts of an AD-PEG stock solution at a concentration of 2.2 mM in water were titrated into the sample cell in 25 separate 10-μL increments. Titrations were performed at 30°C. The measured parameters were $\delta m$, the number of moles of ligand (AD-PEG) added to the sample cell, and $q$, the amount of heat released or absorbed. The Simplex algorithm in the Origin data analysis software was used to determine the following parameters: $K$, the equilibrium binding constant; $n$, the number of available binding sites; and $AH$, the change in enthalpy. A more in depth description of the thermodynamic analysis applied to ITC is provided by Blandamer et al. (21).

Percentage of AD-PEG$_{5000}$ bound after formulation

The small molecule, lactose (Lac), was attached to the end of AD-PEG$_{5000}$ to enable quantification using the Amplex Red Galactose Oxidase Assay Kit (Molecular Probes). Particles were formulated in a total volume of 100 μL water by adding a 50 μL solution containing CDP-Im and AD-PEG$_{5000}$-Lac (1:1 mole ratio of AD-PEG$_{5000}$-Lac: β-CD) in water to a 50 μL solution of siRNA in water. Control formulations were created by mixing CDP-Im and AD-PEG$_{5000}$-Lac without siRNA in 100 μL water. Particles and control formulations were filtered with Biomax (Millipore) centrifugal filtration devices with a 50 kDa MWCO to separate free and bound components. The Amplex Red Galactose Oxidase Assay Kit (Molecular Probes) was used to quantify the amount of AD-PEG$_{5000}$-Lac in the filtrate and retentate of all samples. Concentrations were determined by comparison to a standard curve of AD-PEG$_{5000}$-Lac. The percentage of AD-PEG$_{5000}$-Lac bound to the particles was determined by subtracting the fraction of recovered AD-PEG$_{5000}$-Lac in the filtrate of the
particle samples from the fraction of recovered AD-PEG$_{5000}$-Lac in the filtrate of the control samples.

**Percentage of AD-PEG$_{5000}$-Tf bound after formulation**

Tf-targeted particles were formulated in a total volume of 100 μL water by adding a 50 μL solution of CDP-Im, AD-PEG$_{5000}$, and AD-PEG$_{5000}$-Tf (1:1 mole ratio of AD-PEG$_{5000}$-X:β-CD where AD-PEG$_{5000}$-X was composed of either 1 mol% or 5 mol% AD-PEG$_{5000}$-Tf and the remainder AD-PEG$_{5000}$) in water to a 50 μL solution of siRNA in water. Control formulations were created by mixing CDP-Im, AD-PEG$_{5000}$, and AD-PEG$_{5000}$-Tf without siRNA in 100 μL water. Particles and control formulations were filtered with Nanosep (Millipore) centrifugal filtration devices with a 300 kDa MWCO to separate free and bound components. Total protein content in the filtrate (unbound AD-PEG$_{5000}$-Tf) and retentate (bound AD-PEG$_{5000}$-Tf) was determined using the BioRad DC protein assay. The percentage of AD-PEG$_{5000}$-Tf bound to the particles was determined by subtracting the fraction of recovered AD-PEG$_{5000}$-Tf in the filtrate of the particle samples from the fraction of recovered AD-PEG$_{5000}$-Tf in the filtrate of the control samples.

**Multi-angle light scattering (MALS)**

Particle formulations were loaded into a 10-mL syringe connected to a syringe pump to control the flow rate into a Dawn EOS (Wyatt Technology) multi-angle light scattering instrument. The typical flow rate used was 1 mL min$^{-1}$. Data were fit by the Astra software to the Debye model with a detector fit degree of 2. The $dn/dc$ value for the particles was determined to be 0.14 mL g$^{-1}$, and the mass concentration used in the calculations was determined from the total amount of CDP-Im and nucleic acid incorporated into the particles assuming an individual particle charge ratio of 1.1 (+/−) and complete incorporation of the nucleic acid added during formulation.

**Individual particle stoichiometry**

An estimate for the stoichiometry of each particle (i.e., number of CDP, nucleic acid, AD-PEG, and AD-PEG-Tf molecules) can be calculated from the following equations.

\[
MW_{part} = \#NA \times \#bp \times MW_{bp} + \#CDP \times MW_{CDP} + \#PEG \times MW_{PEG} + \#Tf \times MW_{Tf}
\]  

(1)

\[
CR = \frac{\#CDP}{\#NA \times \#bp}
\]  

(2)

\[
\#PEG = f_{PEG} \times (100\% - \%Tf) \times \#CDP \times PR \times \frac{FR}{CR}
\]  

(3)

\[
\#Tf = f_{Tf} \times \%Tf \times \#CDP \times PR \times \frac{FR}{CR}
\]  

(4)

where \#NA is the number of nucleic acid molecules in the particle, \#CDP is the number of CDP monomers (β-CD) in the particle, \#PEG is the number of AD-PEG molecules in the particle, \#Tf is the number of AD-PEG-Tf molecules in the particle, \(MW_{part}\) is the molecular weight of an individual particle (determined by MALS), \#bp is the number of base pairs per nucleic acid molecule (e.g., 21 for siRNA), \(MW_{bp}\) is the average molecular weight of each nucleic acid base pair (~650 Da as an approximation), \(MW_{CDP}\) is the molecular weight of each CDP monomer (~1,460 Da), \(MW_{PEG}\) is the molecular weight of each AD-PEG molecule (~5,200 Da for AD-PEG$_{5000}$), \(MW_{Tf}\) is the molecular weight of each AD-PEG-Tf molecule (~85,000 Da for AD-PEG$_{5000}$-Tf), \(f_{PEG}\) is the fraction of the AD-PEG molecules that bind to particles during formulation, \(PR\) is the mole ratio of AD-PEG to β-CD during formulation.
FR is the formulation charge ratio (+/−), CR is the charge ratio (+/− = 1.1) of each individual particle, \( f_T \) is the fraction of the AD-PEG-Tf molecules that bind to particles during formulation, and \( \%Tf \) is the mole percent AD-PEG-Tf during formulation.

Salt stability

Particle formulations were diluted to a volume of 1260 μL, placed in a cuvette, and inserted into a ZetaPALS (Brookhaven Instruments Corporation) instrument. Kinetic studies of aggregation were performed by recording the effective diameters at 1 minute intervals after the addition of 1/10 volume 10X PBS to achieve a final concentration of 1X PBS, corresponding to physiological salt concentration.

Erythrocyte aggregation

Erythrocytes were obtained from whole bovine calf blood (Rockland Immunochemicals, Inc.) by multiple rounds of centrifugation at 700xg and 4°C for 10 min followed by removal of the supernatant and resuspension of the pellet of erythrocytes in cold PBS (Cellgro) until the supernatant became clear. Finally, the erythrocytes were resuspended at a concentration of 1% (v/v). The free polycations or formulated particles were added to a 24-well plate and diluted with PBS to a volume of 100 μL. Subsequently, 100 μL of the erythrocyte suspension were added to each well and the plate was incubated for 1 h at 37°C. Images were taken of each well using a CCD-IRIS/RGB (Sony) video camera attached to an Eclipse TE-300 (Nikon) inverted microscope to visually determine the degree of aggregation.

Complement fixation

To test the complement fixation by polycations or CDP-based particles, antibody-sensitized sheep erythrocytes were used in a CH50 assay modified from Plank et al. (11). 25 μL human complement sera (Sigma) in gelatin veronal buffer (Sigma) were added in a 1:1.5 dilution series across a row of wells in a 96-well plate. To this same row of wells were added 25 μL of the desired concentration of polycation in its free form or complexed with calf thymus DNA (CT-DNA). A different concentration of the polycations or particles was added to each row of wells. After a 30-min incubation at 37°C, 1.25×10⁷ antibody-sensitized sheep erythrocytes (Sigma) were added to each well and the plate was incubated with shaking for 1 h at 37°C. Finally, the plate was centrifuged at 2,000 RPM for 10 min, 100 μL of the supernatant from each well was transferred to a new 96-well plate, and the absorbance at 410 nm was determined using a SpectraMax 190 (Molecular Devices) microplate reader. This wavelength corresponds to an absorbance peak for the hemoglobin that is released after lysis of the erythrocytes. The CH50 unit is used to define the serum dilution required to achieve 50% lysis of the antibody-sensitized sheep erythrocytes. If the substance being tested binds complement proteins to an appreciable degree, it will sequester these complement proteins and prevent them from binding to and lysing the erythrocytes. As a result, a lower serum dilution (CH50) will be required to achieve 50% erythrocyte lysis under these conditions. The reported \( \%CH50_{max} \) represents the ratio of the CH50 for the substance being tested to the CH50 determined for the complement sera alone (CH50max).

Cellular uptake

**Method 1: Flow cytometry**—A FACS Calibur (BD Biosciences) flow cytometer was used to detect the uptake of FL-siRNA (fluorescein attached to the 5’ end of the sense strand) delivered with or without the CDP-Im delivery vehicle. HeLa cells were seeded at 2x10⁴ cells per well in 24-well plates 2–3 days prior to transfection and grown in medium supplemented with 10% fetal bovine serum (FBS) and antibiotics (penicillin/streptomycin). The growth medium was removed from each well and replaced with 200 μL Opti-MEM I (Invitrogen) alone, 200 μL Opti-MEM I with 100 nM FL-siRNA, or 200 μL Opti-MEM I with 100 nM FL-
siRNA formulated into CDP-Im particles at a charge ratio of 3 (+−). After incubation for 2 h at 37°C and 5% CO₂, the transfection medium was removed and the cells were trypsinized and resuspended in Hanks Balanced Salt Solution (HBSS) with 1% bovine serum albumin (BSA) and 10 μg mL⁻¹ propidium iodide to detect cell viability.

**Method 2: Confocal microscopy**—HeLa cells were seeded at 2×10⁴ cells per well in a LabTek II Chamber Slide 2 days prior to transfection and grown in medium supplemented with 10% FBS and antibiotics (penicillin/streptomycin). The growth medium was removed from each well and replaced with 200 μL Opti-MEM I containing 100 nM FL-siRNA formulated into CDP-Im particles at a charge ratio of 3 (+−). After incubation for 2 h at 37°C and 5% CO₂, cells were fixed for 15 min at room temperature using 4% paraformaldehyde in PBS. F-actin was stained with rhodamine phalloidin (Invitrogen) according to manufacturer’s instructions. Cells were mounted with Biomeda Gel/Mount according to manufacturer’s instructions and the coverslips were subsequently sealed using nail polish. Fluorescent images were acquired using a Zeiss LSM 510 Meta laser scanning confocal microscope with a 40X water-immersion objective.

**Competitive uptake**

Competitive uptake studies were conducted to determine the impact of free holo-transferrin (holo-Tf) on the relative uptake of transferrin-targeted (containing 1 mol% AD-PEG₅₀₀₀-Tf) or non-targeted particles. By formulating the particles with Cy3-siRNA (Cy3 attached to the 5’ end of the sense strand), a Tecan SPECTRAFluorPlus plate reader could be used to measure the total cell-associated fluorescence after transfection. Cells were seeded at 2×10⁴ cells per well in 24-well plates 2–3 days prior to transfection and grown in medium supplemented with 10% FBS and antibiotics (penicillin/streptomycin). The growth medium was removed from each well and replaced with 200 μL Opti-MEM I containing 100 nM Cy3-siRNA formulated in particles. After incubation for 30 min at 37°C and 5% CO₂, the transfection medium was removed and the cells were lysed in 100 μL cell lysate buffer (Promega). Total fluorescence in the 100 μL lysate per well was measured with the SPECTRAFluorPlus plate reader and the number of siRNA molecules was estimated from a standard curve of Cy3-siRNA. Cells in two wells that were not transfected were trypsinized and counted to provide an estimate for the average number of cells per well.

**Avidity effects**

**Method 1: Competitive cell-surface transferrin receptor (TfR) binding assay**—Competitive uptake experiments were performed using flow cytometry to detect the uptake of fluorescently labeled holo-Tf. Unlabeled holo-Tf, Tf conjugates (AD-PEG₅₀₀₀-Tf), or Tf-targeted siRNA particles (1 mol% AD-PEG₅₀₀₀-Tf) were used to compete for uptake by the transferrin receptors on the surface of HeLa cells. Cells were seeded at 2×10⁴ cells per well in 24-well plates 2–3 days prior to transfection and grown in medium supplemented with 10% FBS and antibiotics (penicillin/streptomycin). The growth medium was removed from each well and replaced with 200 μL Opti-MEM I containing 1% BSA, 20 nM AlexaFluor488-labeled holo-Tf (AF488-Tf), and the desired unlabeled Tf competitor. After incubation for 1 h at 37°C and 5% CO₂, the transfection medium was removed and the cells were trypsinized and resuspended in Hanks Balanced Salt Solution (HBSS) with 1% bovine serum albumin (BSA) and 10 μg mL⁻¹ propidium iodide to detect cell viability. To enable direct comparison of the effects of avidity, the total amount of Tf was kept constant whether it was in its free form, as AD-PEG₅₀₀₀-Tf, or as AD-PEG₅₀₀₀-Tf on the siRNA particles. The relative uptake under each condition is reported as the ratio of the mean fluorescence of the wells with unlabeled competitor to the mean fluorescence of the wells with AF488-Tf alone.
Method 2: Live-cell binding assay—A live-cell binding assay was used to measure the relative binding of transferrin-targeted siRNA particles. 3×10^5 HeLa cells were resuspended in 100 μL PBS in individual microcentrifuge tubes and cooled on ice. To each microcentrifuge tube were added 100 μL PBS containing PEGylated or Tf-targeted particles formulated with Cy3-labeled siRNA such that the final Cy3-siRNA concentration was 100 nM. After incubating for 30 minutes on ice, the microcentrifuge tubes were centrifuged for 5 minutes at 200xg to pellet the cells. 100 μL of the supernatant from each microcentrifuge tube were added to a well in a black 96-well plate, and the Cy3 fluorescence was measured using a Tecan Safire plate reader. Comparison to a standard curve of Cy3-siRNA particles allowed quantification of the amount of Cy3-siRNA in each well, and the percent bound (fraction associated with the cell pellet) was determined by subtracting the fluorescence remaining in the supernatant from the initial amount added.

Luciferase knockdown after siRNA transfection

Functional efficacy of pDNA and siRNA delivered by CDP-Im particles was demonstrated in HeLa cells by co-transfecting the pGL3-CV vector (Promega) containing the firefly luciferase gene and a non-targeting control siRNA (siCON1) synthesized by Dharmacon or a luciferase-targeting siRNA (siLuc) synthesized by Integrated DNA Technologies. The sequence of the siCON1 siRNA is UAGCGACUAAACACAUCAAUU (sense) and UUGAUGUGUUUAGUCGCUAUU (antisense). The sequence of the siLuc siRNA is GUGCCAGAGUCCUUCGAUAdTdT (sense) and UAUCGAAGGACUCUGGCACdTdT (antisense). The Promega Luciferase Assay System was then used to quantify the relative luciferase expression in cells that had been transfected with 1 μg pGL3-CV alone, 1 μg pGL3-CV and 100 nM siCON1, or 1 μg pGL3-CV and 100 nM siLuc. HeLa cells were seeded at 2×10^4 cells per well in 24-well plates 2–3 days prior to transfection and grown in medium supplemented with 10% FBS and antibiotics (penicillin/streptomycin). CDP-Im particles were formulated to contain 1 μg pGL3-CV vector and 100 nM siRNA in 200 μL Opti-MEM I. The growth medium was removed from each well and replaced with 200 μL Opti-MEM I containing the formulated particles. After incubation for 5 h at 37°C and 5% CO₂, 800 μL complete growth medium was added to each well. 48 h later, the cells were lysed in 100 μL 1X Luciferase Cell Culture Lysis Reagent (Promega). 10 μL of the cell lysate were added to 90 μL of the luciferase substrate, and bioluminescence was measured using a MonoLight (Pharmingen) luminometer. 5 μL of the cell lysate were used in a BioRad DC protein assay to determine the protein concentration in each lysate sample. Luciferase activities are reported as relative light units per mg protein.

RESULTS AND DISCUSSION

Particle formation requires a slight excess of positive charge and protects siRNA from degradation in serum

Results from an electrophoretic mobility shift assay (EMSA) demonstrate that siRNA particles completely form at charge ratios (+/−) greater than ~1 to 1.5 (Figure 2A). At sufficiently high charge ratios, the band corresponding to the free nucleic acid becomes undetectable since the nucleic acid remains associated with the particles that have greatly reduced electrophoretic mobility. To determine what portion of the polycations (CDP) added during formulation actually are incorporated into the particles, the free polycations were separated from the particles after formulation. Regardless of the formulation charge ratio up to 30 (+/−), the charge ratio of individual particles remains ~1 (+/−) (Figure 2B). This is consistent with the results shown in Figure 2A where charge ratios slightly greater than 1 were required to achieve complete particle formation. A nuclease stability assay was conducted to determine if the formation of particles could protect the nucleic acid payload from degradation by nucleases present in serum. While naked siRNA degrades rapidly in serum, siRNA within particles is...
protected from significant degradation even after 4 hours in 50% mouse serum (Figure 3). Additionally, the data given in Figure 3 show (i) there is essentially complete encapsulation of the siRNA by the particles, and (ii) when the particles exposed to serum are disrupted with SDS, the nucleic acids released are still intact siRNA duplexes.

**Formulation conditions affect particle size and zeta potential**

Transmission electron microscopy (TEM) and atomic force microscopy (AFM) were used to visualize the siRNA particles formulated at a charge ratio of 3 (+/−). The images in Figure 4 demonstrate that the siRNA particles assume a roughly spherical shape, but the unPEGylated particles display more variability in size and adopt a slightly oblong shape relative to the PEGylated particles when they are visualized by AFM on the mica surface. While a large fraction of the unPEGylated particles (0.1 g L−1 siRNA) have diameters that exceed 100 nm, PEGylated particles (0.5 g L−1 siRNA) formulated with a 1:1 mole ratio of AD-PEG5000:β-CD consistently have diameters <100 nm and are approximately 60–80 nm.

To further investigate the effects of formulation conditions, dynamic light scattering was used to measure the effective hydrodynamic diameter and zeta potential of the particles. Consistent with the TEM and AFM images, the results shown in Figure 5A reveal that the nucleic acid concentration during formulation affects the size of the particles. Particles formulated with siRNA, pDNA, and calf thymus (CT-DNA) show nearly identical trends of increased size with higher nucleic acid concentration. However, particles that are formulated in the presence of AD-PEG5000 (PEGylated particles formed by the pre-complexation method) do not exhibit such a dependence on formulation conditions (Figure 5B). These properties allow the delivery vehicles to be fine-tuned with respect to size by altering the formulation conditions accordingly (i.e., PEGylation through the pre- or post-complexation method). The zeta potential of unPEGylated particles ranges from 10 to 30 mV for similarly sized particles. This positive zeta potential implies that the charge ratio of the individual particles is slightly greater than 1 (+/−). The AD-PEG5000 conjugates can be further modified to contain targeting ligands on the distal end of the PEG chain. For example, transferrin can be conjugated to the AD-PEG5000 molecules to yield AD-PEG5000-Tf (19). Because the transferrin protein is negatively charged, inclusion of AD-PEG5000-Tf molecules during particle formulation reduces the zeta potential of siRNA particles in a concentration-dependent manner (Figure 6). Bellocq et al. reported a similar trend using particles made with pDNA (19).

**AD-PEG conjugates bind to the surface of particles through inclusion complex formation**

An important property of the cyclodextrin-containing polycations is their ability to form inclusion complexes with hydrophobic molecules. This provides the opportunity for modular attachment of different stabilizing molecules or targeting ligands through coupling to an adamantane (AD) molecule that forms inclusion complexes with the β-cyclodextrin molecules. Isothermal titration calorimetry was used to investigate the thermodynamics of the interaction between AD-PEG molecules and CDP-Im either in its free form or within siRNA-containing particles (Table 1). Figure 7 shows representative ITC data plots for binding between AD-PEG5000 and CDP-Im formulated with siRNA at a charge ratio of 3 (+/−), CDP-Im formulated with siRNA at a charge ratio of 10 (+/−), and CDP-Im alone. As the charge ratio increases, the measured binding parameters approach those of free CDP-Im as expected. Given the previous results showing the actual particle charge ratio is slightly greater than 1 (+/−), this is consistent with the presence of excess free CDP-Im at charge ratios >1.

The value of $n$ represents the fraction of the β-CD molecules available for inclusion complex formation with the AD-PEG molecules. Each β-CD cup can interact with a single adamantane molecule, as previous reports have shown a 1:1 binding stoichiometry between adamantane...
Particle molecular weight can be used to estimate individual particle stoichiometry

Determining the individual component stoichiometry of the particles provides important insights into their functional properties. Centrifugal filtration was used to separate unbound components from those bound to the particles. As discussed previously, it was determined that the individual particle charge ratio (i.e., the ratio of positive charges from the CDP-Im to negative charges from the nucleic acid) is slightly greater than 1; we used 1.1 (+/-) for the calculations. The percent of the total AD-PEG5000 or AD-PEG5000-Tf added to the formulation mixture that remains free was determined by quantifying the AD-PEG5000 (experiment actually used AD-PEG5000-Lac and we assumed that the value for AD-PEG5000 would be approximately the same) or AD-PEG5000-Tf recovered in the filtrate versus the retentate after centrifugal filtration. The results of these measurements indicated that approximately 3% of the total AD-PEG5000 and 10% of the total AD-PEG5000-Tf added during formulation remained associated with the particles. The greater degree of binding measured for the AD-PEG5000-Tf conjugates may be partly due to charge interactions between the negatively charged transferrin proteins and the positively charged particles. The final piece of data needed to estimate the individual particle stoichiometry is the molecular weight of the particles. This was determined using multi-angle light scattering. The results in Figure 8 show that the molecular weight of particles formulated with siRNA, pDNA, or calf thymus (CT-DNA) scales approximately as \( r^3 \), where \( r \) is the radius of the particle determined by DLS. This similarity between all three types of particles is consistent with the trends observed in Figure 5, further supporting the interesting result that formulation with a variety of nucleic acids leads to particles with similar physical properties. An unPEGylated particle with a diameter of 70 nm is expected to have a molecular weight around \( 1 \times 10^8 \) g mol\(^{-1} \) from data given in Figure 8. Inserting this molecular weight and an individual particle charge ratio of 1.1 (+/-) into Equations 1 and 2 yields 48,800 CDP monomers (or 9,750 CDP chains with a degree of polymerization of 5) and 2,110 siRNA molecules (with 21 bp per siRNA) per particle. For the sake of calculation, we can then use this value for \#CDP to estimate the number of AD-PEG5000 and AD-PEG5000-Tf molecules per particle using Equations 3 and 4. For example, a 70-nm siRNA particle with a molecular weight of \( 1.3 \times 10^8 \) g mol\(^{-1} \) (accounting for the added mass from the PEG conjugates) formulated at a charge ratio of 3 (+/-) with 1 mol\% AD-PEG5000-Tf is calculated to contain 9,750 CDP chains, 2,110 siRNA molecules, 3,950 AD-PEG5000 molecules, and 133 AD-PEG5000-Tf molecules. The corresponding surface density for this number of AD-PEG5000 chains on a 70-nm particle is \( \sim 43 \) pmol cm\(^{-2} \) or 0.26 chains nm\(^{-2} \). The calculated average

molecules and \( \beta \)-CD (22). While binding between individual \( \beta \)-cyclodextrins and AD-PEG5000 conjugates gives the expected \( n \) value of \( \sim 1 \), binding between CDP-Im and AD-PEG5000 exhibits an \( n \) value of \( \sim 0.5 \). When the AD-PEG5000 molecule containing a 5000-Da PEG chain binds to a \( \beta \)-CD cup on the CDP-Im polymer, it likely provides steric hindrance that impedes binding between other AD-PEG5000 molecules and nearby \( \beta \)-CD molecules. Support for this hypothesis comes from the observation that the \( n \) value for binding between CDP-Im and AD-PEG5000 molecules containing short 500-Da PEG chains is 0.92 \( \pm \) 0.05, likely indicating that the shorter PEG chains do not interfere to as great an extent with the binding of AD-PEG5000 to neighboring \( \beta \)-CD molecules.

Another interesting pattern is observed with the \( \Delta H \) values, representing the change in enthalpy that results from binding between an AD-PEG molecule and a \( \beta \)-CD cup. These values are all negative, indicating that energy is released upon binding due to the favorable interaction between the hydrophobic adamantane and the \( \beta \)-CD cup. Notably, \( \Delta H \) is more negative for AD-PEG5000 binding to the siRNA particles than it is for AD-PEG5000 binding to free CDP-Im. We hypothesize that this increased stabilization energy, in addition to the inclusion complex formation, is a result of favorable interactions between the PEG chains themselves when they are grouped together on the surface of an siRNA particle.
distance between PEG₅₀₀₀₀ molecules at this surface density is ~2.0 nm, while the Flory radius is ~6 nm. Since the distance between PEG₅₀₀₀₀ molecules is much less than the Flory radius, the PEG₅₀₀₀₀ chains are expected to interact laterally and extend out from the surface in a dense brush layer with an estimated thickness of ~12.5 nm. Hansen et al. further examined the brush scaling laws for polyethylene glycols and predicted that PEG₅₀₀₀₀ solutions must have monomer volume fractions, ϕ, greater than 0.07–0.09 to be in the brush regime (23). This is satisfied when the PEG₅₀₀₀₀ surface density exceeds ~26–28 pmol cm⁻², again indicating that the PEG₅₀₀₀₀ chains on the particles are in the brush regime.

**PEGylation provides steric stabilization to the particles and reduces nonspecific interactions**

DLS-based kinetic studies of aggregation were performed to determine whether the inclusion of AD-PEG conjugates could help to stabilize the particles against aggregation at physiological salt concentrations. First, the ratio of AD-PEG₅₀₀₀₀:β-CD (mol:mol) was varied from 0 to 2 to investigate how the surface density of AD-PEG₅₀₀₀₀ affects the steric stability of siRNA particles formulated through the post-complexation method (Figure 9A). Particles formulated with AD-PEG₅₀₀₀₀:β-CD (mol:mol) ratios >1 do not exhibit observable aggregation after 15 minutes in 1X PBS. At ratios <1, aggregation increases as the ratio of AD-PEG₅₀₀₀₀:β-CD (mol:mol) decreases. These results with siRNA-containing particles are consistent with those observed by Pun et al. using particles made with pDNA (24). The length of the PEG molecule in the AD-PEG conjugate also impacts its ability to confer steric stabilization to the particles. As shown in Figure 9B, the stabilization effects increase with the length of the PEG chain, with AD-PEG₅₀₀ : AD-PEG₅₀₀₀₀:β-CD = 1) only slightly slowing the aggregation while AD-PEG₅₀₀₀₀ (AD-PEG₅₀₀₀₀:β-CD = 1) prevents detectable aggregation up to 15 minutes after salt addition. If the AD-PEG₅₀₀₀₀ chains, like the AD-PEG₅₀₀₀₀ chains, also achieve a surface density of ~43 pmol cm⁻² (AD-PEG₅₀₀₀₀:β-CD = 1), then the average distance between PEG₅₀₀₀₀ chains remains ~2.0 nm. However, since this is not less than the Flory radius for a PEG₅₀₀₀₀ molecule (~1.5 nm), the PEG₅₀₀₀₀ molecules are not expected to form the brush-like layer on the particle surface that is needed for steric stabilization. Furthermore, modification of up to 1 mol% of the AD-PEG₅₀₀₀₀ chains with Tf (AD-PEG₅₀₀₀₀-Tf) leads to minimal perturbations in the salt stability of the particles. However, at 5 mol% AD-PEG₅₀₀₀₀-Tf, gradual particle aggregation becomes apparent during the 15-minute incubation in 1X PBS.

Besides providing steric stabilization to the particles, PEGylation can help to reduce nonspecific interactions. Specifically, experiments were performed to study the interaction between the polycations (or particles) and erythrocytes (Figure 10). Significant erythrocyte binding will lead to aggregation that can be observed by visual inspection using a light microscope. While the free CDP or CDP-Im showed negligible aggregation at 0.2 g L⁻¹, some aggregation was observed as the concentration increased to 2 g L⁻¹ (Figure 10A–D). Erythrocyte aggregation was also measured after incubation with siRNA particles that were formulated with CDP-Im and a 1:1 molar ratio of AD-PEG₅₀₀₀₀:CDP-Im (Figure 10E). The results demonstrate that PEGylated particles do not lead to any observable aggregation at a total CDP-Im concentration of 0.2 g L⁻¹, corresponding to the expected concentration after systemic delivery in vivo (7).

**PEGylated particles show minimal complement fixation**

Complement fixation by polyethylenimine and polylysine was compared to that of CDP or CDP-Im. The CDP and CDP-Im molecules do not show as much complement fixation as PEI (branched or linear) or a 36-mer of polylysine, but they do exhibit higher complement fixation than a 5-mer of polylysine (Figure 11A). This is consistent with the notion that polycation length and charge density can augment complement activation (11).
Because complement fixation was observed at the physiologically relevant concentration of 0.2 g L\(^{-1}\), corresponding to the typical concentration of polycations in the bloodstream after delivery of nucleic acids at a dose of 2.5 mg kg\(^{-1}\) (a typical dose used for \textit{in vivo} siRNA delivery (7)), experiments were performed to test particles formulated with calf thymus DNA and stabilized by PEGylation (Figure 11B). Notably, these formulations showed minimal complement fixation at polymer concentrations of 0.2 g L\(^{-1}\).

**Particles achieve intracellular delivery of siRNA \textit{in vitro}\**

The uptake of particles containing fluorescently labeled siRNA was assessed using flow cytometry and confocal fluorescence microscopy. While naked siRNAs do not achieve measurable levels of cellular uptake, formulation into particles with CDP-Im dramatically increases the amount of cell-associated siRNA as measured by flow cytometry (Figure 12A). To confirm that the siRNA was being delivered to the interior of cells, confocal fluorescence microscopy was used to visualize cells transfected with particles containing fluorescently labeled siRNA (Figure 12B). The internalized particles exhibited a punctate staining and were eventually observed to accumulate in the perinuclear region.

**Targeting ligands enhance cellular uptake of PEGylated particles**

To verify that attachment of AD-PEG\(_{5000}\)-Tf can lead to uptake through transferrin receptor (TfR)-mediated endocytosis, the uptake of stabilized (PEGylated) particles was measured in the presence or absence of free holo-Tf. While the uptake of PEGylated particles without AD-PEG\(_{5000}\)-Tf was not affected by the presence of free holo-Tf, the uptake of Tf-targeted particles was reduced as a result of competition with free holo-Tf (Figure 13). Because the particles can be internalized simultaneously by numerous mechanisms including simple pinocytosis, there is still significant uptake even without TfR-mediated internalization under these conditions.

**Targeted particles exhibit increased affinity through avidity effects**

If multiple receptor/ligand interactions can occur simultaneously, then the effective affinity of the binding interaction can be enhanced through avidity effects. Antibodies or divalent antibody fragments are excellent examples of molecules whose binding affinities are enhanced through avidity effects. Their divalent interactions allow single antibodies to bind two separate receptors, leading to a stronger apparent affinity than exhibited by the monovalent antibody fragment (25). Targeted delivery vehicles that contain multiple targeting ligands on the surface should also display these avidity effects if multiple targeting ligands can simultaneously interact with the receptors. A typical cancer cell may contain thousands of receptors on its surface (26,27), and the Tf-targeted delivery vehicles can contain tens or even hundreds of Tf ligands (depending on the percent of the AD-PEG\(_{5000}\) molecules with Tf molecules attached to the distal end of the flexible PEG\(_{5000}\) chains) decorating each particle surface. This arrangement should enable multiple Tf molecules to bind to TfR on the surface of the cells. To test whether these avidity effects increase the apparent affinity of the Tf-targeted particles for the TfR on the cell surface, a competitive uptake assay was performed using flow cytometry. The results shown in Figure 14A demonstrate that the Tf-targeted particles possess enhanced affinity for the TfR relative to individual AD-PEG\(_{5000}\)-Tf molecules. Additionally, particles without the Tf targeting ligand had a minimal impact on the uptake of the fluorescently labeled holo-Tf. To determine how targeting ligand density affects particle binding to cell-surface TfR, particles were incubated with HeLa cells in PBS at 4°C to measure the amount of binding in the absence of internalization. The results shown in Figure 14B show that Tf targeting increases the amount of cell-associated particles under these conditions, and the amount of binding increases with the targeting ligand density.
Particles deliver functional pDNA and siRNA to cells in vitro

In addition to achieving intracellular delivery of the nucleic acid-containing particles, the particles need to release their nucleic acid payload intracellularly to allow it to function. Co-delivery of a luciferase-expressing plasmid and either a control or luciferase-targeting siRNA was used to demonstrate the ability of the particles to deliver functional pDNA and siRNA. The luciferase activity in cell lysates was quantified using a luminometer, and relative light units (RLU) were normalized to total cellular protein levels. As shown in Figure 15, cells that received CDP-Im particles containing the plasmid and siRNA against luciferase (siLuc) had luciferase activity that was ~50% lower than cells that received CDP-Im particles with either the plasmid alone or the plasmid plus a control siRNA (siCON1).

Nanoparticles are multifunctional, integrated systems for nucleic acid delivery

The results presented here highlight the importance of creating a nanoparticle that consists of multiple components that function together as a system, and control over size, surface modification, payload protection, and targeting ligand to payload ratio are key parameters to consider when designing nucleic acid delivery vehicles for in vivo systemic use. These parameters also represent some of the major advantages of nanoparticle composites for delivery of nucleic acids instead of using carrier-free delivery methods. Nucleic acid delivery vehicles can help reduce renal clearance while adding features such as stabilization against nuclease degradation, cell-specific targeting, and large payload delivery. These features make them well-suited for the systemic delivery of nucleic acids in general, and we have shown that the system investigated here can deliver pDNA, siRNA, and DNazymes in vitro and in vivo (7, 17,19,20).

The capability to fine-tune the delivery vehicle’s properties combined with an understanding of how those properties affect its function in biological systems represent two key factors necessary for optimization of nucleic acid delivery vehicles. This study demonstrates the importance of a rational approach in delivery vehicle design and lays a foundation for further in vivo studies to understand the relationships between the properties of nanoparticle delivery systems and their biological function. The approach to designing nanoparticle delivery vehicles that is outlined here can be used for other synthetic materials and is thus not limited to the cyclodextrin polymer-based system illustrated.

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References


Figure 1.
Formation of nucleic acid-containing particles using CDP-Im. (A) Schematic of the chemical structure of CDP-Im. (B) Schematic of particle assembly.
Figure 2.
Effect of formulation charge ratio (+/-). (A) Electrophoretic gel mobility shift assay demonstrating the effect of formulation charge ratio on siRNA particle formation. (B) Individual particle charge ratio as a function of formulation charge ratio.
Figure 3.
Nuclease stability of siRNA encapsulated within particles. For the t = 4 lanes, naked siRNA or siRNA within CDP-Im particles (3 (+/-)) was incubated in 50% mouse serum for 4 h at 37° C and 5% CO₂. For the t = 0 lanes, serum was added to an equivalent amount of naked siRNA or siRNA within CDP-Im particles immediately before loading into the gel. Addition of 1% SDS was used to displace the siRNA from the particles to visualize the amount of intact siRNA remaining. The first lane demonstrates that the upper bands are nonspecific bands resulting from the interaction between SDS, serum, and the ethidium bromide stain, while the lower bands correspond to the free siRNA.
Figure 4.
Transmission electron microscopy (left panels) and atomic force microscopy (center and right panels) images of (A) unPEGylated and (B) PEGylated siRNA particles formulated at a charge ratio of 3 (+/−). Scale bar = 100 nm (left panels) and 200 nm (center and right panels).
Figure 5.
Effect of nucleic acid concentration ([NA]) during formulation on the size of (A) unPEGylated or (B) PEGylated particles. Particles were formulated at a charge ratio of 3 (+/-) using CDP-Im and either siRNA, pDNA, or CT-DNA (calf thymus DNA). PEGylated particles were formulated by adding a 1:1 mole ratio of AD-PEG5000β-CD. Particle effective diameter was determined using dynamic light scattering. Squares = CDP-Im/siRNA particles, circles = CDP-Im/pDNA particles, diamonds = CDP-Im/CT-DNA particles.
Figure 6.
Particle zeta potential as a function of AD-PEG$_{5000}$-Tf ligand concentration during formulation. Particles were formulated at a charge ratio of 3 (+/−) using CDP-Im and siRNA, and the AD-PEG$_{5000}$ or AD-PEG$_{5000}$-Tf molecules were added after particle formation (post-complexation). The total number of moles of AD-PEG$_{5000}$-X (AD-PEG$_{5000}$ and AD-PEG$_{5000}$-Tf) was equal to the number of moles of β-CD, and the mixture of AD-PEG$_{5000}$ and AD-PEG$_{5000}$-Tf is defined by the % AD-PEG$_{5000}$-Tf.
Figure 7.
Isothermal titration calorimetry (ITC) plots characterizing the binding between AD-PEG5000 molecules and free CDP-Im or siRNA particles. (A) CDP-Im/siRNA particle (3 (+/−)) and AD-PEG5000- (B) CDP-Im/siRNA particle (10 (+/−)) and AD-PEG5000- (C) CDP-Im and AD-PEG5000.
Figure 8.
Relationship between particle size and molecular weight (MW) as determined by dynamic and multi-angle light scattering. Particles were formulated at a charge ratio of 3 (+/−) using CDP-Im and either siRNA, pDNA, or CT-DNA. Effective diameters were measured using dynamic light scattering, and molecular weights were determined using multi-angle light scattering. Squares = CDP-Im/siRNA particles, circles = CDP-Im/pDNA particles, diamonds = CDP-Im/CT-DNA particles, solid line = \( r^3 \) scaling dependence of the MW of particles starting with a MW of \( 7 \times 10^7 \) g mol\(^{-1} \) for a 60-nm particle.
Figure 9.
Aggregation of siRNA particles in physiological salt solutions. 140 μL of a 10X PBS solution were added to 1260 μL of the particles in water after 5 minutes, and dynamic light scattering was used to follow the formation of aggregates with time. (A) Effect of the ratio of AD-PEG<sub>5000</sub>:β-CD on particle stability. CDP-Im/siRNA (3 (+/−)) particles were formulated without AD-PEG<sub>5000</sub> (black squares) or through the post-complexation method with an AD-PEG<sub>5000</sub>:β-CD mole ratio of 0.25:1 (black triangles), 0.5:1 (inverted black triangles), 0.75:1 (black diamonds), 1:1 (black circles), or 2:1 (black stars). (B) Effect of PEG chain length, adamantane conjugation, and Tf targeting ligand density on particle stability. CDP-Im/siRNA (3 (+/−)) particles were formulated without AD-PEG<sub>5000</sub> (black squares), with a PEG<sub>5000</sub> (no
adamantane):β-CD mole ratio of 1:1 (open inverted triangles), with an AD-PEG$_{5000}$-β-CD mole ratio of 1:1 (black triangles), with an AD-PEG$_{5000}$:β-CD mole ratio of 1:1 (black circles), or with a 1:1 mole ratio of AD-PEG$_{5000}$-X:β-CD where AD-PEG$_{5000}$-X was composed of 0.1 wt % AD-PEG$_{5000}$-Tf (dark gray circles), 0.1 mol% AD-PEG$_{5000}$-Tf (gray circles), 1 mol% AD-PEG$_{5000}$-Tf (light gray circles), or 5 mol% AD-PEG$_{5000}$-Tf (open circles) and the remainder AD-PEG$_{5000}$. 
Figure 10.
Erythrocyte aggregation. (A) 0.2 g L$^{-1}$ CDP, (B) 2 g L$^{-1}$ CDP, (C) 0.2 g L$^{-1}$ CDP-Im, (D) 2 g L$^{-1}$ CDP-Im, (E) CDP-Im/siRNA (3 (+/−)) particles at 0.2 g L$^{-1}$ CDP-Im formulated with a 1:1 mole ratio of AD-PEG$_{5000}$:β-CD, (F) PBS alone. Scale bar = 20 μm.
Figure 11.
Complement fixation. (A) Complement fixation by free polycations. Asterisks = branched PEI, x = linear PEI, black triangles = pentalysine, inverted black triangles = polylysine (36-mer), black squares = CDP, black circles = CDP-Im. (B) Complement fixation by CDP/CT-DNA (3 (+/−)) particles formulated with a 1:1 mole ratio of AD-PEG5000β-CD (black squares). The curves for CDP (open squares), CDP-Im (open circles), and pentalysine (open triangles) are shown again for comparison.
Figure 12.
Uptake of CDP-Im particles containing fluorescein (FL)-labeled siRNA by HeLa cells. (A) Histogram of cell-associated fluorescence measured by flow cytometry. The left-most peaks correspond to the overlapping peaks for HeLa cells incubated with either Opti-MEM I alone or 100 nM naked FL-siRNA, while the right-most peak represents the cell-associated fluorescence after transfection with CDP-Im/FL-siRNA particles. (B) Confocal fluorescence microscopy image of HeLa cells after transfection with CDP-Im/FL-siRNA particles (green) and rhodamine phalloidin staining of F-actin (red). Scale bar = 50 μm.
Figure 13.
Uptake of PEGylated and Tf-targeted particles in the presence of holo-Tf competitor. Particles were formulated at a charge ratio of 3 (+/−) using CDP·Im and siRNA. PEGylated particles (PEGpart) were formulated with a 1:1 mole ratio of AD-PEG\textsubscript{5000}·β-CD and Tf-targeted particles (Tfpart) were formulated with a 1:1 mole ratio of AD-PEG\textsubscript{5000}·X·β-CD where AD-PEG\textsubscript{5000}·X was composed of 99 mol% AD-PEG\textsubscript{5000} and 1 mol% AD-PEG\textsubscript{5000}·Tf. Particles containing 100 nM siRNA were added to HeLa cells in 200 μL Opti-MEM I in the absence or presence of a 25X (moles holo-Tf: moles AD-PEG\textsubscript{5000}·Tf) excess of holo-Tf competitor.

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Figure 14.
Effect of transferrin targeting ligand density on relative binding affinity. (A) Competitive TfR binding by free holo-Tf (circles), free AD-PEG<sub>5000</sub>-Tf (triangles), Tf-targeted CDP-Im/siRNA (3 (+/−), 1 mol% AD-PEG<sub>5000</sub>-Tf) particles (squares), and PEGylated CDP-Im/siRNA (3 (+/−)) particles (diamonds) in the presence of 20 nM AlexaFluor488-labeled holo-Tf. As a control, the PEGylated particles were formulated identically to the Tf-targeted particles at each concentration except without the addition of AD-PEG<sub>5000</sub>-Tf during formulation. (B) Live-cell binding assay. Particles were formulated at a charge ratio of 3 (+/−) using CDP-Im and Cy3-labeled siRNA. PEGylated particles (PEGpart) were formulated with a 1:1 mole ratio of AD-PEG<sub>5000</sub>−β-CD and Tf-targeted particles (Tfpart) were formulated with a 1:1 mole ratio
of AD-PEG5000-X:β-CD where AD-PEG5000-X was composed of the stated % AD-PEG5000-Tf and the remainder AD-PEG5000. Particles containing 100 nM Cy3-siRNA were added to HeLa cells in 200 μL PBS and incubated on ice for 30 minutes. The “percent bound” represents the fraction of particles associated with the cell pellet after centrifugation.
Figure 15.
Luciferase expression 48 h after co-transfection of HeLa cells with particles containing pDNA and siRNA. Particles were formulated at a charge ratio of 3 (+/-) by combining CDP-Im with pGL3-CV (pGL3 Alone), pGL3-CV and a control siRNA (pGL3+siCON1), or pGL3-CV and an siRNA against luciferase (pGL3+siLuc).
Table 1
Measured ITC parameters for the binding between AD-PEG<sub>5000</sub> and β-CD alone, polycation alone (CDP-Im), or CDP-Im/siRNA particles formulated at charge ratios from 3 (+/−) to 15 (+/−). For comparison, literature values are provided for the binding between β-CD alone and adamantane carboxylate (22).

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