Improved siRNA delivery efficiency via solvent-induced condensation of micellar nanoparticles

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Abstract

Efficient delivery of short interfering RNA (siRNA) remains one of the primary challenges of RNA interference therapy. Polyethylene glycol (PEG)ylated polycationic carriers have been widely used for the condensation of DNA and RNA molecules into complex-core micelles. The PEG corona of such nanoparticles can significantly improve their colloidal stability in serum, but PEGylation of the carriers also reduces their condensation capacity, hindering the generation of micellar particles with sufficient complex stability. This presents a particularly significant challenge for packaging siRNA into complex micelles, as it has a much smaller size and more rigid chain structure than DNA plasmids. Here, we report a new method to enhance the condensation of siRNA with PEGylated linear polyethylenimine using organic solvent and to prepare smaller siRNA nanoparticles with a more extended PEG corona and consequently higher stability. As a proof of principle, we have demonstrated the improved gene knockdown efficiency resulting from the reduced siRNA micelle size in mice livers following intravenous administration.

Supplementary material for this article is available online

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(Some figures may appear in colour only in the online journal)

Introduction

RNA interference (RNAi) has been demonstrated to be a potent and highly specific post-translational gene regulation...
process [1]. As it specifically targets the gene of interest through the use of short interfering RNA (siRNA), RNAi holds great potential as a therapeutic agent for the treatment of numerous disorders [2, 3]. However, delivery of naked siRNA molecules via intravenous injection has failed to yield significant gene knockdown due to their poor pharmacokinetic profile, resulting from a high susceptibility to nuclease degradation and rapid renal clearance [3]. Thus, it is a considerable challenge to balance PEI-siRNA complex stability with colloidal stability and thus, it is a considerable challenge to balance PEI-siRNA complex stability with colloidal stability and ease of formulation [4]. Polycationic nanoparticles have gained significant attention owing to their versatility and ease of formulation [5]. Polyethylenimine (PEI) is the most commonly used polycationic carrier, due to its high buffering capacity that facilitates endosomal escape of nanoparticles and siRNA into the cytoplasm [6]. However, PEI/siRNA nanoparticles typically carry positive residual charges on their surface, which lead to significant aggregation in the presence of serum proteins, thereby greatly reducing their efficacy in vivo [7]. Furthermore, they often elicit a high degree of inflammatory responses and toxicity, and are prone to clearance by macrophages following systemic administration, particularly for branched PEI [8, 9].

Decorating the particle surface with hydrophilic polyethylene glycol (PEG) has often been employed as an effective strategy to alleviate aggregation, reduce opsonization and inflammation response, and prolong nanoparticle circulation time [10]. PEGylation can be achieved by grafting PEG chains onto the polycation backbone. The resulting PEGylated polycations retain their ability to complex with siRNA, forming micellar nanoparticles with a polycation/siRNA core and a PEG corona, where the steric shielding effect of the corona has been demonstrated to lessen aggregation of PEI-g-PEG/siRNA nanoparticles [11, 12]. However, whereas a higher degree of PEG grafting density favors the colloidal stability and biocompatibility of the micelles, it reduces the RNA-condensation capacity of the PEI-g-PEG carrier [13]. Thus, it is a considerable challenge to balance PEI-g-PEG/siRNA complex stability with colloidal stability and compatibility.

Experimental methods

Preparation of linear polyethylenimine (IPEI)-g-PEG/siRNA micelles

The siRNA (1.33 μg, Qiagen, Valencia, CA) was first dissolved in 50 μl of deionized water or a 7:3 (v/v) dimethylformamide (DMF)–water mixture and then added to an equal volume of IPEI-g-PEG polymer solution at an N/P ratio of 20 prepared in the same mixing solvent. The mixture was vortexed and then incubated for 30 min at room temperature before further characterization. Crosslinked micelles were prepared and purified according to a protocol that we established previously [14]. Micelle purification and characterization methods are provided in the online supplementary information.

In vitro cell uptake and gene silencing of IPEI-g-PEG/siRNA micelles

HepG2 cells were maintained in Dulbecco’s Modified Eagle’s Medium supplemented with 10% fetal bovine serum and 100 U ml⁻¹ Penicillin/100 μg ml⁻¹ Streptomycin at 37 °C and 5% CO₂. At 24 h prior to the experiment, cells were seeded in 24-well plates at a density of 5 × 10⁴ cells/well. For cell-uptake studies, micelles were prepared with Alexa Fluor 488-modified siRNA (Qiagen, Valencia, CA). An aliquot of 50 μl micelles equivalent to 100 nM siRNA was added to each well followed by 4 h incubation at 37 °C, after which the cells were washed with phosphate-buffered saline (PBS), trypsinized, and fixed with 2% paraformaldehyde. Fluorescence associated with individual cells was analyzed with a BD FACSCalibur flow cytometer (BD Biosciences, San Jose, CA) fitted with a 488 nm excitation source and detected using a 515–545 nm filter. A minimum of 10,000 events per sample was collected for analysis. Gene-silencing studies were performed using a previously reported protocol [15].

In vivo gene knockdown efficiency of siRNA micelles via intravenous administration

Animal studies were conducted under a protocol that was approved by the Johns Hopkins School of Medicine Institutional Animal Care and Use Committee (IACUC # RA09A447). A liver-specific gene-knockdown model was developed by modification of previously published protocols [16, 17]. Wistar rats (female, 6–8 weeks, 200–300 g) were transfected with a firefly luciferase plasmid DNA (20 μg DNA in PBS with a volume corresponding to 9 vol/wt% of the rat’s body weight) via hydrodynamic infusion administered through the tail vein over 15 s according to a published procedure [18]. After 5 d, micelles containing 80 μg siRNA in 1 ml PBS were injected via the tail vein. At 24 and 48 h, rats were anesthetized and given 1 ml of D-luciferin solution (i.p. 30 mg ml⁻¹). Each rat was then imaged on an IVIS Spectrum Imaging System. The bioluminescence signal was collected for 1 min, and the level of luciferase expression was expressed as the total photon count per section in the region of interest and was normalized to PBS control to determine knockdown efficiency.

Statistical analysis

All data were expressed as mean ± SD unless otherwise noted. Statistical comparisons were carried out using a one-way analysis of variance followed by Tukey’s post-hoc test for groups with equal variance or Games–Howell test for groups with unequal variance (SPSS software, version 21,
IBM Inc., Armonk, NY). All data were considered to be significant at $p < 0.05$.

**Computational methods**

**Modeling of IPEI-g-PEG/siRNA micelles**

Molecular dynamics (MD) simulations were performed using coarse-grained models. IPEI-g-PEG was represented as a bead–spring polymer model with bead size 7.35 Å and charge density 35%. The siRNA molecule was coarse-grained as a 24-bead rigid body using the VMD Shape-Based Coarse-Graining tool [19, 20], based on a 22-bp RNA molecule isolated from the Protein Data Bank file 2FS8 [21]. To increase the computational efficiency, we scaled down each coarse-grained model to one fourth of its original length. The solvent was simulated implicitly using a Langevin thermostat, and different solvent compositions were represented through variation of the attractive strength of a Lennard–Jones potential. Monovalent counterions were included to maintain charge neutrality, and electrostatic interactions were computed using the Ewald method. To accelerate the dissociation and reformation of aggregates (to improve sampling of aggregate conformations), we employed the parallel tempering method [22], in which 24 copies of the same system were simulated in parallel, at closely spaced temperatures. This approach exploits the larger degree of fluctuations at higher temperatures to provide pathways that permit the simulation at the original temperature to transition between different states of low free energy that are separated by free-energy barriers.

**Results and discussion**

**siRNA condensation and size control**

Our new approach for condensing siRNA employs a PEGylated linear PEI (IPEI) carrier, IPEI-g-PEG, which was synthesized by grafting PEG ($M_n = 10\, \text{kJDa}$) to the backbone of PEI ($M_n = 17\, \text{kDa}$; see the online supplementary information for copolymer synthesis and characterization). We prepared an IPEI-g-PEG copolymer with an average of 4.6 PEG grafts per PEI, and identified a minimal N/P ratio of 20 for complete siRNA condensation by agarose gel retardation assay (figure S1 is available at stacks.iop.org/NANO/28/204002/mmedia). For all following experiments, we prepared siRNA micelles by mixing equal volumes of 200 $\mu$g ml$^{-1}$ of IPEI-g-PEG solution and 20 $\mu$g ml$^{-1}$ of siRNA solution (corresponding to an N/P ratio of 20) at room temperature. The intensity-averaged diameter of these IPEI-g-PEG/siRNA micelles prepared in water was $117 \pm 2.3\, \text{nm}$, as measured by dynamic light scattering (DLS; figures 1(A) and (F)). On the other hand, IPEI/siRNA nanoparticles exhibited a slightly smaller diameter of $96.8 \pm 10.3\, \text{nm}$ (figures 1(A) and (E)). Whereas this size difference may be attributed to the PEG corona, it is nevertheless remarkable given our observations for plasmid DNA, where condensation with block copolymers significantly reduced the size of the micelles compared to condensation with polyelectrolytes alone [14].

To understand this apparent discrepancy, we performed MD simulations of complexation between siRNA and IPEI-g-PEG copolymer. We emphasize that both the time scale and the number of particles involved in the complexation processes make coarse-graining (as described in the online supplementary material) imperative. Whereas this approach omits atomistic details, prior work has shown it to be highly suitable for providing meaningful insight into the underlying mechanisms [14, 23]. In addition to the electrostatic interactions between siRNA and IPEI (taken into account via Ewald summation [24–26]), and the IPEI–solvent and PEG–solvent interactions [14], we also incorporated the interaction between IPEI and PEG arising from direct hydrogen bonding and from hydrogen bond bridges with water molecules, as IPEI is known to form strong inter- and intra-molecular hydrogen bonds in water [27]. Given that PEG not only has a molecular structure similar to IPEI, but also has a hydrogen bond acceptor in its repeat unit, we adopted the same effective interaction strength for PEG–IPEI as for IPEI–IPEI [27]. To compare to the intensity-averaged distribution of the hydrodynamic diameter measured by DLS, we report the z-averaged [28] radius of gyration of the IPEI-g-PEG/siRNA nanoparticles calculated in simulation (figure 1(B)). The modeling results indeed matched the experimental observation that the complexation of siRNA with IPEI-g-PEG copolymer results in a slight increase in particle size. This increase did not arise merely from the physical extent of the PEG corona: the average number of siRNA particles per micelle increased from 7.1 $\pm$ 1.3 to 9.4 $\pm$ 1.3. We hypothesized that this size increase can be attributed to the IPEI–PEG interaction. To test this conjecture, we performed a second set of simulations of the complexation of siRNA with IPEI-g-PEG copolymer, in which we artificially decreased the attractive interaction between IPEI and PEG (online supplementary figure S4). We indeed observed that a weakening of the PEG–IPEI interaction significantly reduces the particle size.

Prompted by this observation, we aimed to condense IPEI-g-PEG/siRNA micelles into smaller nanoparticles by disrupting the interaction between IPEI and PEG mediated by hydrogen bonding with water molecules. We added DMF, a water-miscible solvent that has been shown to effectively reduce intermolecular hydrogen bonding in other polymeric micelle systems [29], to siRNA and IPEI-g-PEG solutions prior to micelle assembly. In a 7:3 (v/v) DMF–water mixture, we indeed observed a far smaller average particle size of $44.2 \pm 6.6\, \text{nm}$ (figures 1(C) and (G)). This size reduction of micelles in response to a decrease in solvent polarity was confirmed by MD simulations (figure 1(D)). Arguably, the polarity reduction also leads to a decrease in siRNA solubility [30], which in turn could have resulted in stronger condensation and a growth in aggregate size [14]. We conclude that this effect is overshadowed by the reduction of PEG–IPEI and IPEI–IPEI hydrogen bonding which in turn permits the PEG corona to provide steric shielding.
Preservation of nanoparticle size following solvent removal

Whereas DMF as a co-solvent can significantly reduce particle size, it needs to be removed prior to nanoparticle transfection experiments in vitro or in vivo. However, DMF removal by dialysis resulted in nanoparticle swelling (data not shown). To preserve the nanoparticle size in aqueous media upon DMF removal, we employed a reversible disulfide crosslinking scheme previously tested in DNA/polymer nanoparticles [14]. We introduced thiol groups to the IPEI block and used the purified thiolated IPEI-g-PEG to condense siRNA using the protocol described above. Crosslinking was initiated by aerial oxidation over a 48 h incubation at room temperature followed by dialysis to remove the DMF. TEM micrographs (figures 2(A) and (B)), confirm that no appreciable change in particle size or morphology occurred.
We anticipated a smaller average particle size to result in more efficient cellular uptake and transfection. To confirm this, we first measured the cellular uptake efficiency and gene knockdown efficiency in vitro. We also prepared 117 nm micelles that were crosslinked similarly to the 44 nm micelles to exclude differences resulting from the crosslinking. The crosslinked 44 nm micelles displayed 30% and 57% increase of uptake in a hepatocellular carcinoma cell line (HepG2) compared to the crosslinked and uncrosslinked 117 nm micelles, respectively (figure 3(A)). In vitro gene knockdown experiments showed that both the crosslinked 117 and 44 nm micelles achieved about 60% decrease in targeted protein expression, an efficiency similar to that of IPEI-siRNA nanoparticles. The uncrosslinked micelles only showed 30% knockdown efficiency, likely due to their limited stability in cell culture media (figure 3(B)). In addition, all tested particles maintained a high cell metabolic activity (>80% compared to untreated cells, online supplementary figure S3). As an in vivo proof of concept, we assessed the effect of reduced nanoparticle size on gene-knockdown efficiency in the rat liver following intravenous administration (figure 3(C)). We first established high luciferase expression in the liver via hydrodynamic infusion of luciferase plasmid DNA [31]. After the transgene expression level stabilized at 5 d post-infusion, IPEI-g-PEG/siRNA micelles or IPEI/siRNA nanoparticles containing 80 μg luciferase siRNA were administered via tail vein injection. The 44 nm micelles displayed 85% and 70% efficiencies of transgene knockdown in the liver at 24 h and 48 h after injection, respectively. In contrast, the 117 nm micelles only showed ~30% knockdown at both 24 h and 48 h (p < 0.05). Control nanoparticles delivering a non-targeting sequence also did not show any reduction in luciferase expression. This significant increase in gene knockdown efficiency for the smaller nanoparticles may be attributed to improved nanoparticle deposition and increased cellular uptake of the smaller nanoparticles. On the other hand, the

The PEG corona of IPEI-g-PEG/siRNA micelles significantly reduced the nanoparticle zeta potential from +30 mV to about +5 mV when condensed in water. Condensation in DMF-water (7:3, v/v) mixture followed by crosslinking and removal of solvent, reduced the zeta potential even further to ~8 mV (online supplementary figure S2), likely due to the increase of the PEG density at the surface and reduction of imines on the complex core that reacted with the crosslinkers. The near-neutral surface charge on these micelles enhances their colloidal stability in physiological media through suppression of serum protein-mediated agglomeration. Indeed, these crosslinked micelles showed no significant change in particle size in 10% serum (figures 2(D) and (E)). Owing to the covalent crosslinks, these micelles also exhibited high complex stability; they did not show any size change when incubated in the presence of 150 mM sodium chloride (figure 2(F)). On the other hand, uncrosslinked IPEI-g-PEG/siRNA micelles were destabilized within 30 min of salt challenge.

**siRNA delivery and knockdown efficiency**

We anticipated a smaller average particle size to result in more efficient cellular uptake and transfection. To confirm this, we first measured the cellular uptake efficiency and gene knockdown efficiency in vitro. We also prepared 117 nm micelles that were crosslinked similarly to the 44 nm micelles to exclude differences resulting from the crosslinking. The crosslinked 44 nm micelles displayed 30% and 57% increase of uptake in a hepatocellular carcinoma cell line (HepG2) compared to the crosslinked and uncrosslinked 117 nm micelles, respectively (figure 3(A)). In vitro gene knockdown experiments showed that both the crosslinked 117 and 44 nm micelles achieved about 60% decrease in targeted protein expression, an efficiency similar to that of IPEI/siRNA nanoparticles. The uncrosslinked micelles only showed 30% knockdown efficiency, likely due to their limited stability in cell culture media (figure 3(B)). In addition, all tested particles maintained a high cell metabolic activity (>80% compared to untreated cells, online supplementary figure S3). As an in vivo proof of concept, we assessed the effect of reduced nanoparticle size on gene-knockdown efficiency in the rat liver following intravenous administration (figure 3(C)). We first established high luciferase expression in the liver via hydrodynamic infusion of luciferase plasmid DNA [31]. After the transgene expression level stabilized at 5 d post-infusion, IPEI-g-PEG/siRNA micelles or IPEI/siRNA nanoparticles containing 80 μg luciferase siRNA were administered via tail vein injection. The 44 nm micelles displayed 85% and 70% efficiencies of transgene knockdown in the liver at 24 h and 48 h after injection, respectively. In contrast, the 117 nm micelles only showed ~30% knockdown at both 24 h and 48 h (p < 0.05). Control nanoparticles delivering a non-targeting sequence also did not show any reduction in luciferase expression. This significant increase in gene knockdown efficiency for the smaller nanoparticles may be attributed to improved nanoparticle deposition and increased cellular uptake of the smaller nanoparticles. On the other hand, the
uncrosslinked micelles and lPEI/siRNA nanoparticles did not show significant knockdown at 24 h and yielded a low level (≈15%) of transgene knockdown at 48 h after injection, suggesting that nanoparticle stability in medium is crucial to siRNA delivery in vivo. It is important to note that the gene knockdown activity mediated by the 44 nm siRNA nanoparticles is among the highest obtained via intravenous injection at such a relatively low siRNA dose (≈0.4 mg kg⁻¹ body weight), without employing any active targeting strategy [32–34].

**Conclusion**

In conclusion, this study provides the first evidence of solvent-assisted condensation of lPEI-g-PEG/siRNA micelles to decrease nanoparticle size. By reducing solvent polarity, we decreased the average size of lPEI-g-PEG/siRNA micelles from 117 nm in water to 44 nm. Through MD simulation we revealed the role of solvent quality and PEI–PEG hydrogen bonding in the assembly of lPEI-g-PEG/siRNA micelles. The micelle size was preserved after organic solvent removal by means of reversible disulfide crosslinking; we confirmed that the micelles maintained their size in water and physiological media. More importantly, our results have demonstrated size-dependent in vivo transfection efficiency following intravenous injection of the siRNA micelles in rats. The gene knockdown efficiency in rat liver achieved by the smaller siRNA micelles was significantly higher than for the larger micelles prepared from the same copolymer carrier. The condensation technique introduced here allows a simple and effective way to reduce siRNA particle size and provides a model platform for further study of the effect of particle size on in vivo cellular uptake, knockdown efficiency, biodistribution, and pharmacokinetics. As such, it can form the starting point for the development of an effective delivery system that harnesses the therapeutic potential of siRNA.

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References

[26] Sanders L K et al 2007 Control of electrostatic interactions between F-actin and genetically modified lysozyme in aqueous media Proc. Natl Acad. Sci. USA 104 15994–9