Cationic vesicles for efficient shRNA transfection in the MCF-7 breast cancer cell line

Introduction: Novel and safe delivery solutions for RNAi therapeutics are essential to obtain the full potential of cancer gene therapy.

Methods: In this study, cationic vesicular nanocarrier was applied for delivering lnc urothelial carcinoma-associated 1 (Inc UCA1) shRNA expression vector to MCF-7 cells. The physicochemical characteristics, cytotoxicity, and transfection efficiency of cationic vesicles prepared from various molar ratios of amphiphilic surfactant Tween 80 (T), squalene (S), cationic charge lipid didodecyldimethylammonium bromide, and polyethylenimine were investigated. The particle sizes of the vesicles in the nanosize range were determined by dynamic light scattering and transmission electron microscopy.

Results: Gel protection assay with agarose gel electrophoresis showed cationic vesicles can protect the shRNA plasmid from DNase 1 enzyme. 3-(4,5-Dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H tetrazolium, inner salt result showed no significant cytotoxicity was caused in MCF-7 cancer cell line by (T:S) polyethylenimine cationic vesicles. 3-(4,5-Dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H tetrazolium, inner salt assay, fluorescence microscope images, and flow cytometry analyses confirmed that (T:S)P \_1,040 \_g/mL with 4.3 \_g/mL of PEI vesicles provided effective transfection without significant cytotoxicity. Furthermore, we found efficient UCA1 shRNA transfection and significant (P<0.05) cell cycle arrest and apoptosis in MCF-7 cancer cells.

Conclusion: The novel nonviral vesicular nanocarrier, (T:S)P \_1,040 \_μM with 4.3 \_g/mL of PEI, might be safe and efficient for cancer gene therapy and can be used in further in vitro and in vivo studies.

Keywords: cationic vesicles, shRNA, UCA1, apoptosis, gene delivery

Introduction

Recent nanotechnology advances empower drug delivery in medical sciences. Nonviral vectors are appropriate candidates for efficient and safe gene delivery. Vesicles which comprise nonionic surfactants and excipients, known as niosomes, are safe and not immunogen.1 Some excipients similar to cholesterol or squalene can be applied for stabilizing the vesicular structure and others such as cationic lipids or polymers can be used to achieve appropriate properties for the nucleic acid transfection.2,4 Niosomes potentially ameliorate drug delivery and bioavailability and show enhancement effects on factors such as stability, solubility, entry and controlled release of the drugs, simultaneously, decreasing immunogenicity.5,6 As a preliminary study of ocular gene therapy, Puras et al7 developed a cationic niosome containing a plasmid that efficiently transfected into the retina of rats without significant angiogenesis.

Gene therapy researchers have provided enormous hope to treat a wide variety of cancers, infections, respiratory, and nervous diseases.8-11 Overcoming the systemic
delivery of RNAi is a hard challenge to produce gene drugs. RNAi by shRNA can be stable even for 2 years in comparison with siRNA that degrades 99% in 2 days. The synthesis cost, fast systemic clearance, weak cell entrance, and lack of specificity for tissue targeting are considered the main barriers for potential clinical applications of siRNA. Here we present shRNA with survivin promoter to achieve cancer cell targeting. Several tumor-specific promoters such as survivin were evaluated for targeted cancer gene therapy. Survivin is an antiapoptotic gene overexpressed in a wide range of human cancers with a low or no expression in normal tissues.

Recently, genes in regulatory noncoding RNAs have been considered for gene therapy studies. Urothelial carcinoma-associated 1 (UCA1) long noncoding RNA was discovered for the first time in bladder cancer. This long noncoding RNA forms a complex with heterologous nuclear ribonucleoprotein; its expression is mainly restricted in the nucleus and it is also involved in pre-mRNA processing. Although there is little information about the molecular mechanisms underlying UCA1 biological activity, especially its antiapoptotic effects, UCA1 has known effects on cell proliferation, motility, invasion of tumors, and drug resistance in different cancers.

Cationic lipids are efficient excipients for plasmid transfection. Plasmid transfection improvement has been reported in the lungs of BALB/c mice and in rat retina, in an animal model survey. In another study, a cationic niosome formed efficient complexes for codelivery of an anticancer phytochemical and a resistance-related gene siRNA in MCF-7 breast cancer cells and nude mice. Opanasopit et al reported high transfection efficiency and low cytotoxicity of cationic niosomes prepared in vitro by span 20, cholesterol, and novel spermine-based lipids as cationic counterparts. More recently, Tagalakis et al developed a bioengineered nanovesicle platform by synthesis and nanoformulation of cationic peptides, which provide the appropriate positive zeta potentials and targeting simultaneously. These giant unilamellar vesicles make stable nanosize complexes with safe and efficient nucleic acid transfection in vitro and in vivo.

The adenoviruses-based gene delivery systems are more efficient for plasmid delivery relative to branched polyethylenimine (PEI)-based polyplexes, but PEI-based nanoformulations for modification in binding, uptake, endosomal escape, and cell nuclear entrance could enhance gene delivery efficiency close to that of adenoviruses for future potential application of nonviral drug delivery systems.

In this study, we describe the application of several cationic vesicle formulations comprising a cationic lipid (didodecyldimethylammonium bromide [DDAB]), a cationic polymer (PEI), squalene, and Tween 80 (polysorbate 80), with the ability of transfection of a large plasmid (6.5 kb) containing the UCA1 shRNA. Additionally, we explore the antiapoptotic role of UCA1 in the MCF-7 breast cancer cell line. There are few studies focusing on the antiapoptotic effect of UCA1 in human breast cancer. Here, the UCA1 IncRNA shRNA expression vector with survivin promoter was constructed and delivered by vesicular nanocarriers to MCF-7 cells.

Materials and methods

Materials

DDAB, squalene, Tween 80, and PEI with an average molecular weight of 25,000 (PEI-25k) were purchased from Sigma-Aldrich Co. (St Louis, MO, USA). Culture medium (RPMI-1640) and FBS were purchased from Thermo Fisher Scientific (Waltham, MA, USA). Dulbecco’s PBS was purchased from Thermo Fisher Scientific. Ultrapure water was obtained from EMD Millipore RIOs™. Annexin V Apoptosis Detection kit was purchased from BioLegend (Shanghai, China). MCF-7 (human breast adenocarcinoma cell line) was obtained from Pasteur Institute (Tehran, Iran). 3-(4,5-Dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H tetrazolium, inner salt (MTS) was purchased from Promega Corporation (Fitchburg, WI, USA). Reverse transcription-PCR (RT-PCR) Master Mix Reagents kit and cDNA Synthesis kit were purchased from Fermentas. Other chemicals were of reagent grade and used without further purification.

Construction of UCA1 shRNA expression vector

Using bioinformatics tools, the core promoter sequence of survivin (Sur-p; 269 bp) was selected and cytomegalovirus (CMV) enhancer was placed upstream the survivin core promoter to improve promoter activity. Thereafter, sequence alignment was performed in CLUSTAL omega to find conserved sequences between UCA1 gene variants and a specific shRNA was designed via InvivoGen-siRNA Wizard software and placed downstream of enhancer/promoter. Finally, the 772 bp nucleotide fragment was synthesized by Shine Gene Molecular Biotech Company and subsequently cloned in the pRNAT-U6.1/Neo expression vector (6,380 bp; GenScript Corp., Piscataway, NJ, USA) to construct the pRNAT-CMVhancer-Sur-p-sh (UCA-1) vector. Double digestion analysis was used to verify substitution of the U6 promoter into the CMVhancer-Sur-p. Plasmid DNA was extracted from Escherichia coli transformants using Fastfilter Endo-free Plasmid Midi kit (Omega Bio-Tek Inc., Doraville, GA, USA).
Preparation of the complexes

Briefly, DDAB cationic lipid was dispersed in PBS solution containing Tween 80 emulsifier and squalene. First, Tween 80 and squalene were dissolved in chloroform and this emulsion was then shaken for 4 hours at 37°C for thin film formation. Then, PBS containing DDAB and PEI was added to the thin film and then the emulsion was vortexed for 30 seconds and incubated for 40 minutes at 37°C until it became clear and transparent. To get different molar and charge ratios, lipoplexes were incubated for 30 minutes at room temperature before use to enhance electrostatic interactions.

Binding, DNase I protection, and SDS-induced release of DNA

Vectors in a concentration of 0.03 μg DNA/μL were subjected to electrophoresis on an ethidium bromide-containing gel (1% agarose). Subsequently, bands were photographed with a Vilber (Vilber Lourmat, Marne-la-Vallée, France) E-BOX. In the protection study, 1 U of DNase and 1.2 μg DNA (Sigma-Aldrich) were incubated (37°C) with the vectors and complexes for 30 minutes. Then, a 2% sodium dodecyl sulfate (SDS) solution was added as a DNA release reagent. Samples were subjected to agarose gel electrophoresis and compared to untreated DNA.

Physical properties of particles: size, morphology and zeta potential

Dynamic light scattering (DLS) assay was performed to determine the physical properties of the cationic vesicles. The zeta potential, size, and polydispersity index (PDI) of the vesicles in PBS (pH 7.4) were analyzed by DLS (Zetasizer Nano ZS; Malvern Instruments, Malvern, UK) using an argon laser beam at 633 nm and a 90° scattering angle.

Transmission electron microscopy (TEM) and ultraviolet (UV)–visible spectroscopy

TEM (EM10C; Carl Zeiss Meditec AG, Jena, Germany) was used to observe the morphology, size, and polydispersity of complexes. Complexes were sonicated for 10 minutes in a bath ultrasonicator. Thereafter, a small drop of the sample solution was deposited on a copper grid covered by 0.2% polyvinyl formal (Vinylec K). The grid was allowed to dry at room temperature. Prior to observation, complexes were negatively stained with a solution of 2% uranyl acetate. The UV-visible spectra were obtained using a Cary 60 UV-visible Spectrophotometer (Agilent Technologies, Santa Clara, CA, USA) in an absorbance range between 200 and 700 nm.

Human monocyte isolation

Informed consent letters were obtained from the blood donors, and the toxicity evaluation study in peripheral blood mononuclear cells (PBMCs) was approved by Golestan University of Medical Sciences Ethics Committee. Peripheral venous blood samples (10 mL) were taken by venipuncture from two students, and isolation of PBMCs with Lymphodex (Inno-Train 002041600) was performed according to the manufacturer’s protocol. Briefly, 10 mL of peripheral blood was mixed with 1 mL sodium citrate. The mixture of blood and PBS (1:4, v:v) was mildly shaken and the tubes were centrifuged for 3 minutes at 1,000×g at room temperature. The white pellet obtained was cautiously transferred to 4 mL of cooled Lymphodex. Then, again, the tubes were centrifuged for 30 minutes at 1,000×g at room temperature. Aspiration of white rings of lymphocytes was done cautiously with a Pasteur pipette and washing with PBS solution was performed. Then, the pellet was washed three times and centrifuged at 250×g, 175×g, and 150×g for 10 minutes. After the last centrifugation, PBMCs were cultivated in six-well plates in a humidified atmosphere at 37°C, 5% CO₂ for 1–3 hours. The supernatant medium (which contained nonadherent cells) was then removed and the adherent cell layer was washed twice with warm PBS and replaced by fresh DMEM (Thermo Fisher Scientific) with 10% FBS (Thermo Fisher Scientific). The following day, the cells were detached by adding ice-cold 0.02% EDTA/PBS solution for 10 minutes and then firmly tapping the flask to obtain mononuclear cells.

Cell culture

Human breast MCF-7 and MDA-MB-231 cancer cell lines were obtained from the Pasteur Institute. MCF-7, MDA-MB-231, and extracted mononuclear cells were cultured in RPMI 1640 medium (Thermo Fisher Scientific) supplemented with 10% FBS and penicillin–streptomycin solution at 37°C in a humidified atmosphere of 5% CO₂. After third passage, the cells were split and used for further experiments.

Cell toxicity assay

MCF-7 and MDA-MB-231 cells were seeded at 1×10⁴ cells per well in a 96-well plate in a final volume of 200 μL/well at −24 hours before the assay. Cells were exposed to different complexes at various concentrations. At the end of the incubation time (72 hours), 20 μL/well MTS reagent (Promega Corporation) was added to each sample and the plates were incubated for 2.5 hours at 37°C in standard culture conditions. Finally, the absorbances were measured at 490 nm.
using ELISA reader (Awareness Technology ChroMate® Microplate Reader). The estimated percentage of cell viability was compared with the value of the untreated control cells.

Cell transfection
The knockdown of Inc-UCA1 was performed by synthetic shRNA. A day before transfection, cells were seeded in six-well plates at a density of 3×10⁴ cells per well in 2.0 mL RPMI 1640 growth medium with 10% FBS. Prior to transfection, cationic vesicular complexes were prepared as described above and added to the wells to a final volume of 2 mL with serum-free transfection medium. Then, after 5 hours of incubation, the transfection mixture was replaced with fresh medium containing serum and cells were maintained under normal growth conditions. Finally, after 72 hours of transfection, the cells were harvested in Trizol (Tri Reagent; Sigma-Aldrich) for RNA isolation.

RNA extraction and quantitative RT-PCR analysis
The treated and untreated cells were washed with PBS and harvested following 72 hours of incubation. Total RNA was extracted from the cells using Trizol reagent according to the manufacturer’s instruction. RNAs were reverse transcribed using the PrimeScript RT reagent kit (Thermo Fisher Scientific). The quantitative PCR experiments were used for the analysis of UCA1 gene expression using a standard SYBR Green PCR kit (Thermo Fisher Scientific) protocol. PCR reactions were run in triplicate for three independent experiments. The sequences of the primers (Metabion) are as follows: UCA1F, 5’ AGCTAACTGGCACCTGTG 3’, UCA1R, 5’ TGGGGATTACTGGGGTAGGG 3’, and 18SF and 18SR as internal control were 5’ GATACCCGTTGACCCCGATT 3’ and 5’ CCATCCAATCGTTAAGCGCC 3’, respectively.

Cell cycle analysis
Cells were harvested 72 hours after transfection, washed with ice-cold PBS, and fixed with 75% ethanol for 2 hours at 4°C. Cells were washed and suspended in PBS containing 100 μg/mL RNase A (Thermo Fisher Scientific) and then incubated at 37°C for 30 minutes to eliminate the intracellular RNA. Finally, cells were stained with propidium iodide (PI; 10 μg/mL; Sigma-Aldrich) in dark for 15 minutes. The results were assessed by flow cytometer (BD Biosciences, San Jose, CA, USA). The percentages of the cells in G0/G1, S, and G2/M phases were determined and compared with the control group. Each experiment was performed in triplicate.

Apoptosis detection
The apoptotic index of each sample was measured through flow cytometry assays using FITC Annexin V and PI staining kits (BioLegend®, San Diego, CA, USA) according to the manufacturer’s instructions. MCF-7 cells were seeded (0.5×10⁶ cells/well) into six-well plates and allowed to adhere overnight before treatments. After 72 hours trypsin-digested cells were centrifuged at 1,100 rpm for 5 minutes. Cells were washed twice with cold BioLegend’s cell staining buffer and then resuspended in Annexin V binding buffer at a concentration of 5×10⁵ cells/mL. One hundred microliters of the cell suspension was transferred to a 5 mL test tube. Five microliters of fluorescein isothiocyanate Annexin V and 10 μL of PI solution were added. After gentle vortexing, the cells were incubated for 15 minutes at room temperature (25°C) in the dark. Four hundred microliters of Annexin V binding buffer was added to each tube and immediately analyzed using BD accurate C6 flow cytometer. The data were evaluated using a BD Accuri™ C6 software (BD Biosciences).

Evaluation of gene delivery efficiency in vivo
The efficiency of gene delivery of cationic vesicles into the tumor cells was evaluated by pRNAT-U6.1/Neo expression vector that consists of a green fluorescent protein gene under the control of CMV promoter. After 1-week acclimatization of female BALB/c mice (8 weeks old; weighing 20 g), subcutaneous tumors were established by injecting 4T1 (4×10⁶ cells in 100 μL PBS) into the upward side of the right hind leg of mice. When the tumor size reached 100 mm³ in volume, the complex of plasmid with cationic vesicles was injected intratumorally.

Immunohistochemistry of GFP
Cryoprotection method was used for the preparation of the tissue sections. Vector-treated tumor was resected from mice and placed in cold PBS to wash out the blood. Then the tumor was cut into appropriate sizes and frozen. The frozen tissues were cut into thin sections by a cryostat at −20°C and transferred onto slides. After fixation, the tissue sections were incubated in DAPI solution for 10 minutes to dye the nuclei. Finally, the slides were observed by fluorescence microscopy.

Statistical analysis
Statistical analysis was performed using SPSS 19.0 software. Results were expressed as mean±SD. Statistical significance
among groups was determined by one-way ANOVA and Student’s t-test. P-values <0.05 were considered statistically significant.

Results and discussion

UCA1 shRNA expression vector with survivin promoter

Multiple sequence alignment of UCA1 gene variants with CLUSTAL omega was done and the shRNA sequence of 5′-ggtaaATGTATCATCGGCTTAGttcaagagaCTAAGCCGA TGATACATTACctaaataaa-3′ was designed for one conserved region of UCA1 gene sequences. The final sequence (772 bp) was cloned into the BglII and HindIII restriction sites of the pRNAT-U6.1/Neo expression vector and verified by double digestion and DNA sequencing.

DNase I protection of plasmid DNA by vesicular formulations

Technically, agarose gel DNase I protection assay shows the efficient ratios of cationic nanocarrier/plasmid formulations which can protect DNA from degradation. Figure 1 shows that free plasmid DNA was degraded completely when exposed to the DNase I (Lane L). The presence of plasmid bands shows that all the vesicles containing PEI protected the DNA against enzymatic digestion. Incubation with DNase I for 30 minutes at 37°C degraded plasmid DNA in the vesicles which contained only squalene and Tween 80 (520 and 1,040 μM; Lanes E and F, respectively). Additionally, vesicles containing DDAB 42, 62.5, and 84 μM could not protect DNA properly. Therefore, all treatments containing PEI were selected for further experiments in this study.

Physical properties of vesicles: size, morphology, and zeta potential and UV-visible spectra

Evaluation of size, zeta potential, and morphology of the nanocarriers was performed by DLS (Table 1) and TEM (Figure 2). Generally, the DLS and TEM results showed that the cationic vesicles comprising Tween 80, squalene, DDAB, and PEI had two particle forms, micelle like and niosome (polymersome like). Preferably, if there is more than one population of nanoparticles, then determination of hydrodynamic diameter should be considered based on volume (% vs mean size) parameter of DLS, as represented in Table 1. These data are similar to our previous result obtained with monomethoxy poly [ethylene glycol]-oleate vesicles, which showed that lowering the concentration of the blocks leads to increase in vesicle size from micellar (10–30 nm) to polymersomes (usually larger than 100 nm),26 and at higher concentration of blocks, the micellar form is dominant in which the formation of micelles besides larger vesicles such as niosomes or polymersomes (comprising lipid-size surfactants) has been widely reported.27 It seems that thermodynamically, there is a tendency to form two populations of micelles and larger vesicles. Another explanation of formation of small micellar size particles is that the potential molecular condensation of plasmid DNA with cationic counterpart may yield small-sized complexes which are close to viruses and micelles. Similar to our result, Perales et al obtained small transfecting particles, 10–12 nm in diameter, with poly(l-lysine) and described successful in vivo gene transfer by tuning the parameters of the polyplex formation, such as the ionic strength of the solvent.28

Interestingly, the DLS data showed that using a higher concentration of DDAB increased the PDI of the cationic

![Figure 1 DNA protection assay. All lanes (except K) represent DNase I-treated contents. Notes: (A) 50 bp DNA ladder. (B–D) Vesicles containing squalene, Tween 80, and DDAB. (E–F) Vesicles containing only squalene and Tween 80. (G–I) Vesicles containing squalene, Tween 80, DDAB, and PEI. (J) Vesicles containing squalene, Tween 80, and PEI. (K) Control (free DNA without DNase I). (L) Control (free DNA with DNase I). Abbreviations: DDAB, didodecyldimethylammonium bromide; PEl, polyethyleneimine; S, squalene; T, Tween 80.](https://www.dovepress.com/10.2147/ijn.s125791ضا،)
vesicles (PDI of 0.677 for 84 μM compared with 0.244 for 42 μM DDAB), as found in Table 1. This result is in accordance with that of Zabner et al which demonstrated that the population of lipid–DNA complexes has a wide range polydispersity.29

The positive zeta potential of the complexes comprising plasmid DNA and the cationic vesicles (ranged from +3.26 to +7.75 mV; Table 1) was induced by DDAB and PEI in the structure. It seems that their positive charges are adequate for the vesicular stability (no precipitation observed by naked eye after 30 days). In addition, this slightly positive net charge potentially can promote interaction with the negatively charged cell surface and the cell entrance.

The UV-visible spectra were recorded using a spectrophotometer (Cary 60 UV-visible) in an absorbance range between 200 and 700 nm (Figure 3). When we examined the UV-visible spectra of (T:S)\textsubscript{520 μM} :DDAB\textsubscript{84 μM}:PEI, (T:S)\textsubscript{333 μM} :DDAB\textsubscript{84 μM}:PEI, and (T:S)\textsubscript{107.1 μM} :PEI, no change in the absorption spectra were not significant changed and increased from 280 to 285 nm in (T:S)\textsubscript{520 μM} :DDAB\textsubscript{84 μM}:PEI. No change in the absorption spectra probably shows weak molecular interactions between plasmid DNA and vesicles containing DDAB, which is in accordance with the DNase I protection assay results.

**Table 1** Size distribution and zeta potential of cationic vesicles

| Treatments | Particle size (nm) | PDI | Zeta potential (mV) |
|------------|-------------------|-----|---------------------|
| \((T:S)_{520 μM}\) | Micelles: 12.71±3.082 (99.6%) | 0.146 | −4.85 |
| \((T:S)_{520 μM}:DDAB_{84 μM}:PEI\) | Niosome (polymersomes): 107.5±30.37 (0.4%) | 0.244 | +7.75 |
| \((T:S)_{333 μM}:DDAB_{84 μM}:PEI\) | | 0.677 | +7.47 |
| \((T:S)_{107.1 μM}:PEI\) | | 0.330 | +6.16 |

Notes: Size is measured as hydrodynamic diameter. All data are shown as volume % vs mean size±SD. Abbreviations: DDAB, didodecyldimethylammonium bromide; PEI, polyethyleneimine; PDI, polydispersity index; S, squalene; T, Tween 80.

**UCA1 expression knockdown by shRNA**

To evaluate the efficacy of gene silencing with designed shRNA vector, UCA1 gene expression analysis was performed in MCF-7 cells. The expression inhibition rates are shown in Figure 4. Following 72 hours of transfection, UCA1 gene expression level revealed a significant reduction in all treated cells with different nanoformulations compared to the nontreated controls.

From a biological viewpoint, some recent reports demonstrate the negative correlation between UCA1 expression with caspase 3, tumor-suppressive miR-143, and metastasis in the breast cancer cell line.30,31 Han et al showed that increased expression of UCA1 leads to cell proliferation and antiapoptotic effect in colorectal cancer cell line and downregulation of UCA1 causes cancer cell arrest.32 A more recent survey in breast cancer showed that the expression of UCA1 increases and promotes cell proliferation in vitro and in vivo.19

**Transfection efficiency**

Transfection efficiency of 6,512 bp UCA1 shRNA plasmid was evaluated in a wide range of (0.5–10 mM) of three ingredients: surfactant (Tween 80; T), squalene (S), and cationic lipid (didodecyldimethylammonium bromide [DDAB]) at the molar ratio of 1 (S):1 (T):(0.08, 0.12, 0.16) (DDAB). Surfactant concentrations >2.5 mM caused detachment of the adherent MCF-7 cells and were evaluated as toxic for further applications. Therefore, lower concentrations have been considered for the following experiments. Efficient transfection in different S:T:DDAB complexes was
Cationic vesicles for efficient shRNA transfection

of lipoplexes comprising Tween 80 (520 or 1,040 μM), squalene (520 or 1,040 μM), and cationic DDAB (42, 62.5, or 84 μM).

Accordingly, PEI as a cationic polymer that improves plasmid DNA transfection efficiency was applied to the T:S-DDAB lipoplex. To understand the role of DDAB combined with PEI, three different concentrations of DDAB (42, 62.5, and 84 μM; DDAB) were selected for further experiments and evaluations. Moreover, for understanding the effect of the surfactant (T) and the excipient (S), two concentrations of 520 μM:520 μM and 1,040 μM:1,040 μM of T:S with a constant amount of 1.3 μL PEI (1 mg/mL) were selected for transfection of 5×10^5 MCF-7 cells in 300 μL of medium (4.3 μg/mL). Fluorescent illumination (GFP channel) of the transfected cells with different vesicular formulations is shown compared with Lipofectamine™ 3000 as a positive control (Figure 5). For quantification of the transfection, Lipofectamine 3000 has been considered as a reference (100%). The transfection ability and efficiency of (T:S)_520 μM:DDAB_84 μM:PEI, (T:S)_520 μM:DDAB_62.5 μM:PEI, (T:S)_520 μM:DDAB_42 μM:PEI, (T:S)_1,040 μM:PEI, (T:S)_1,040 μM:PEI, and PEI, were 43%, 39%, 30%, 78%, 48%, and 34%, respectively.
Results showed that DDAB concentrations were not significantly related to the transfection efficiency, whereas applying two times the amount of vesicular components, S and T, increased the transfection efficiency from 48% to 78%.

DDAB has been used as a cationic lipid for liposomal plasmid transfection (usually smaller than 4 kD, based on our knowledge) and it potentially stabilizes liposomes by DNA complexation by charge neutralization. But in this work, it seems that DDAB cannot help in the transfection of large plasmid (6.5 kb). Here, low shRNA transfection efficiency of the vesicles containing DDAB can be interpreted with Zabner et al’s explanation. They reported that the vesicles containing DDAB can form aggregations after endocytosis of cationic DNA lipoplexes. They showed that before the entry of DNA into the cell nucleus, lipid and DNA cannot dissociate properly and, in addition, they have weak endosomal escape, which is highly necessary for gene delivery, and even after the direct nuclear injection of a condensed lipid–DNA complex, they proved that DNA could not have proper transcription. Probably, DDAB as a cationic counterpart hindered successful transfection.

On the other hand, after endocytosis, PEI-DNA polyplexes have a great endosomal escape (proton sponge effect) because PEI leads to an influx of protons and water, and therefore, endosome swelling and disruption happens, which consequently releases the endosome content to the cytoplasm.

Our data showed that PEI polyplex with (T:S)$_{0.40}$μM:PEI formulation even had better transfection (78%) relative to the intact PEI (34%) for 6.5 kD UCA1 shRNA plasmid, and UCA1 shRNA transfection efficiencies in different treatments were in this order: (T:S)$_{0.40}$μM:PEI > (T:S)$_{0.20}$μM:PEI > PEI (intact PEI), when constant amount of 1.3 μg (4.3 μg/mL) of PEI was used in all formulations.

**Cancer cell toxicity**

MTS assay results for the MCF-7 cells and normal monocyte cells by different vesicular formulations after 72 hours of
Cationic vesicles for efficient shRNA transfection

Figure 6 Cytotoxicity of cationic vesicles on MCF-7 and MDA-MB-231 cancer cell lines and normal monocyte cells.

Notes: Cells were treated with different concentrations of DDAB, surfactant, and PEI in vesicular formulations and evaluated at 72 hours after the UCA1 shRNA treatment. The surviving cells were indirectly measured by MTS assay. The results are shown as the means±SD of triplicates, as a percentage relative to the control. Student’s t-test was used for analysis of significant differences (*P<0.05, **P<0.01, ***P<0.001, ****P<0.0001). S:T treatment (second column) and nontreated cells (first column) were used as two controls of our experiment.

Abbreviations: DDAB, didodecyldimethylammonium bromide; MTS, 3-(4,5-dimethylthiazol-2-yl)-5-(3-carboxymethoxyphenyl)-2-(4-sulfophenyl)-2H tetrazolium, inner salt; ns, nonsignificant; PEI, polyethyleneimine; S, squalene; T, Tween 80.

UCA1 shRNA treatment are shown in Figure 6. No significant toxicity was observed for the vesicles composed of Tween 80 and squalene; it can be inferred that this composition promotes normal monocyte cell growth compared to nontreated cells. Furthermore, this formulation of Tween 80 and squalene (S:T) vesicle was considered as a control to find potential significant toxicity of other ingredients, PEI and DDAB. Promotional effect of (S:T) vesicles on normal monocyte cells can be attributed to squalene, as a natural lipid, as Puras et al reported similar result with squalene in the...
HEK-293 and ARPE-19 cells as application of niosomes containing squalene in the rat retina was safe and effective. 35

Compared with PEI alone (IC50 about 7 μg/mL), no significant toxicity was observed in different vesicular formulations containing T, S, and a constant amount of PEI (4.3 μg/mL). Cell growth enhancement and reduction in toxicity were observed when the concentrations of Tween 80 and squalene were doubled (1,040 μM of S and T, shown as (T:S)1,040μM in both cell types.

The results showed that DDAB can be relatively toxic for the normal monocyte cells and MCF-7 cells, especially with higher concentration (84 μM), which is not reported previously. Among many cationic liposomes studied, a few formulations have the chance for clinical applications. The toxicity of some cationic lipids is an important problem in the application of vesicles. Many researchers have shown the toxicity of different cationic liposomes,36 but in most cases, the mechanisms of toxicity are not clear. It is possible that toxicity is caused by the interaction of the cationic lipids with cell organelle membranes36 and, additionally, it seems that the role of lipid molecular structure is more important than the zeta potential of the lipoprotein. In this order, Filion and Phillips reported moderate toxicity of DDAB liposomes and showed that liposomes containing DDAB and other cationic lipids with similar positive zeta potentials have different toxicity in macrophage cells.37 Therefore, the details of the molecular mechanisms of DDAB on cell toxicity remain to be cleared.

DDAB potentially helps in the stability of vesicles by increasing the zeta potential. However, we observed that application of DDAB with PEI increased the relative cell toxicity, especially in higher concentrations (>42 μM; Figure 6), and implementation and interpretation should be considered cautiously for further applications.

Compared to other drug delivery systems, there are some nanocarriers such as polyamidoamine dendrimers whose cytotoxicity is more severe than that of PEI at the same concentration. Although it makes their application in the gene delivery system difficult,38 surface-engineered dendrimers modified with lipids, peptides, polymers, nanoparticles, and other cationic moieties show promising potential for nucleic acid delivery.39

Cell cycle measurement

Cell cycle analysis was conducted to investigate the function of UCA1 following shRNA treatment in MCF-7 and MDA-MB-231 cancer cell lines and to find which nanoformulation is more effective. Flow cytometric analysis indicated that G2/M arrest was increased significantly in all UCA1 shRNA complexes formed by cationic vesicles from 20.9%±0.25% for (T:S)520 μM vesicles (as control) to 32.8%±0.55%, 34.2%±0.33%, 45.5%±11.5%, 32.6%±2.6%, and 37.8±2.8 in the (T:S)520 μM:PEI, (T:S)520 μM:D42 μM:PEI, (T:S)1,040 μM:PEI, (T:S)520 μM:PEI, and lipofectamine treatments, respectively (Figure 7). All treatments were composed of polyplex of UCA1 shRNA expression vector, except the control which contained a U6 plasmid or pRNAT-U6.1/Neo expression vector (without UCA1 shRNA).

Formulations with different amounts of DDAB caused significant cell cycle arrest, but compared with vesicles that only have PEI, it seems that DDAB and PEI do not have a synergistic effect. The vesicles containing only DDAB were not able to induce efficient transfection for a large 6.5 kD plasmid used in this study (data not shown).

In addition, probably a twofold increase in Tween 80, as a surfactant, and squalene, as an excipient, with a constant amount of 1.3 μg of PEI (4.3 μg/mL) improved the efficiency of vesicle formulation for G2/M cell cycle arrest induction in MCF-7 cells. This result is in accordance with more transfection efficiency of (T:S)1,040 μM:PEI relative to (T:S)520 μM:PEI based on fluorescence microscopy imaging and flow cytometry data; it can be interpreted that Tween 80 and squalene can help PEI in plasmid transfection in MCF-7 cells. Additionally, a slight decrease in cell toxicity of MCF-7 cells and normal monocytes was observed for (T:S)1,040 μM:PEI relative to (T:S)520 μM:PEI. Also, our data are in accordance with Li et al’s result which showed UCA1 overexpression reduced MCF-7 cell cycle arrest after tamoxifen treatment.40

Apoptosis in MCF-7 cancer cells

Significant apoptosis was observed following shRNA UCA1 treatment relative to (T:S)520 μM treatment as a control (Student’s t-test analysis, P>0.05; Figure 8). The MCF-7 and MDA-MB-231 cancer cells were exposed (72 hours) to shRNA UCA1 complexed with different cationic vesicle formulations. In this study, total early and late apoptosis were used for the comparisons. Data showed that UCA1 shRNA polyplexes prepared by (T:S)520 μM:D42 μM:PEI, (T:S)1,040 μM:PEI, and (T:S)520 μM:PEI cationic vesicles induced significant apoptosis in MCF-7 cells (P>0.05) relative to the void control U6 plasmid polyplex formed by (T:S)520 μM vesicles.

Huang et al showed G1 cell cycle arrest and increase in P27 protein with UCA1-siRNA in MCF-7 breast cancer cell line. They implied UCA1 RNA–heterologous nuclear ribonucleoprotein complex will competitively reduce the active...
hnRNA, which subsequently decreases translocation of P27 mRNA from the nucleus to the cytoplasm of cells.41

From the nanostructural point of view, our data showed that formulations comprising a higher amount of DDAB (84 μM) caused significant apoptosis. Similar to the cell cycle result, DDAB and PEI do not have a synergistic effect and the amount of 1.3 μg of PEI (4.3 μg/mL) will be sufficient to transfect 6.5 kD UCA1 shRNA plasmid into MCF-7 cancer cells.

Additionally, it seems that twofold increase of Tween 80 and squalene, with a constant amount of PEI, slightly increased the induction of apoptosis (47.3±5.4 for (T:S)1,040 μM:PEI relative to 41.5±0.9 for (T:S)520 μM:PEI vesicles; Figure 8), while toxicity was reduced in MCF-7 cancer cells.

**Efficiency of nanoparticle gene delivery**

The efficiency of gene delivery into the tumor cells was investigated by examining the expression of GFP in tumor tissue 72 hours after injection of the complex of plasmid with (T:S)1,040 μM cationic vesicles. The mice were sacrificed and tumor tissue was harvested and subjected to immunohistochemical staining to track the fluorescence in the tissue sections. Fluorescence microscopy revealed fluorescing cells in the tumor tissue. It confirmed that cationic vesicles...
efficiently transferred the expression vector into the tumor cells (Figure 9).

**Conclusion**

We demonstrated that the downregulation of UCA1 leads to significant cell cycle arrest and apoptosis in MCF-7 cancer cells, which are in accordance with the previous findings of this noncoding RNA. Although the molecular mechanism of UCA1 in cell biology is not fully understood and needs further evaluations, it seems that downregulation of this lncRNA can be an appropriate approach for gene therapy against human breast cancer.

Among 100 different cationic lipids, only a few are appropriate for in vivo applications. Although many attempts were done to understand the physical and biological properties of cationic lipid–DNA for nucleic acid transfection, still mechanisms of the complexation remain to be cleared and need further evaluations. Nonionic surfactant Tween 80 has been commonly used in a range of pharmaceutical products. The combination of Tween 80 and squalene is somehow similar to the formulation of MF59*-adjuvanted influenza vaccine (by Novartis) which is tested in >30 countries and millions of people without significant safety and immunogenicity concerns.

The cationic PEI-based vesicular nucleic acid nanoformulations provided effective transfection with a low level of cytotoxicity. Additionally, significant apoptotic effect of UCA1 shRNA was observed in MCF-7 cancer cells and it

**Figure 8** (Continued)
showed that UCA1 RNAi can be potentially considered as an appropriate approach for breast cancer gene therapy.

The antiapoptotic effects of other regulatory noncoding genes remain to be investigated. Furthermore, other novel cationic polymers such as recombinant cationic polypeptides can be safer and more efficient alternative for cationic counterparts in future gene therapy surveys.

Here, we showed that (T:S)₁₀₄₀ μM combination can compensate the toxicity of PEI (in a sufficient concentration for large plasmid transfection) to MCF-7 and

Figure 8 Apoptosis induction by shRNA UCA1 complexed with different cationic vesicles in MCF-7 (A and B) and MDA-MB-231 (C and D) cancer cell lines. Notes: (A and C) FITC Annexin V and PI staining assay results. (B and D) Early and late apoptosis were used together to show total apoptosis. The (T:S)₁₀₄₀ μM lipoplex was used as a control, which only had a U6 plasmid without shRNA UCA1 transfecting vector. The results are shown as the mean±SD of three independent experiments. Student’s t-test was used for the analysis of significant differences (P<0.05, n=3). Significance is *P<0.5 and **P<0.01.
Abbreviations: DDAB, didodecyldimethylammonium bromide; PEI, polyethyleneimine; PI, propidium iodide; S, squalene; T, Tween 80.

Figure 9 Immunohistochemical staining of breast tumor from mice. Notes: Complex of plasmid with (T:S)₁₀₄₀ μM cationic vesicles were injected in to the tumor. (A) Nuclei. (B) Cells expressing GFP. (C) Merged. Abbreviations: S, squalene; T, Tween 80.
MDA-MB-231 cancer cells. In summary, the vesicular nanocarrier, (T:S)_{1,040 μμ}, with PEI, might be a useful tool for safe and efficient RNAi-based cancer gene therapy.

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Author contributions
VEM designed the study and conceived the experiments with the help of BJ and analyzed the results and prepared the manuscript with the help of PM. PM conducted the experiments. MS and AS helped in spectrophotometer experiments and their analysis. All authors contributed toward data analysis, drafting and critically revising the paper, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

Disclosure
Mahrokh Sharbatkhari is employed by Arya Tina Gene Bio-pharmaceutical Company (Gorgan, Iran); the authors report no other conflicts of interest in this work.

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