Calcium based siRNA coating: a novel approach for knockdown of HER2 gene in MCF-7 cells using gold nanoparticles

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Abstract: Surface functionalization of nanoparticles (NPs) for therapeutic siRNA delivery into cancer cells has gained interest. The present study was designed for surface functionalization of gold nanoparticles (AuNPs) for efficient siRNA delivery and knockdown in cancer cells. In order to achieve this objective, AuNPs were coated with HER2-siRNA in the presence of 11-mercaptopoundecanoic acid (11-MUA), calcium chloride (CaCl2) and polyethyleneimine (PEI) in alternate charge bearing successive layers. MCF-7 cells were cultured and transfected with fabricated assembly of AuNPs. Cytotoxicity analysis revealed that the half inhibitory concentration (IC50) for the formulation was 45.35 nM. Total RNA was isolated from transfected cells, reverse transcribed into complementary DNA (cDNA) and real-time polymerase chain reaction (RT-PCR) was performed. The RT-PCR based delta-delta Ct analysis in treated cells revealed a significant 18.94 times decrease (p<0.001) in the expression of HER2 gene standardized with ACTB housekeeping gene as compared to untreated cells, which makes this formulation a potent approach for siRNA delivery and gene knockdown.

Key words: Gold nanoparticles; HER2; siRNA silencing; MCF-7.

Introduction

Cancer is one of the major causes of death worldwide. Globally, every 1 in 6 deaths is related to cancer and the number is still expected to increase (1). In cancer, cells of the body start dividing uncontrollably and this abnormal division is triggered by mutation in either proto-oncogenes or tumor suppressor genes, resulting in the loss of cell cycle regulation and increased proliferation (2). Traditional strategies for cancer treatment include surgery in conjunction with chemo and/or radiotherapy. Success ratio of these traditional therapeutic methods is not very high and requires cancer to be diagnosed earlier. Gene therapy provides an excellent way to target cancer, especially via gene silencing of up-regulated oncogenes (3). RNA interference (RNAi) is being used to modulate gene expression by utilizing the small fragments of double stranded RNA (dsRNA) molecules including short hairpin RNA (shRNA), endogenous microRNA (miRNA), and small interfering RNA (siRNA). The most utilized one is siRNA due to the ease of synthesis and no requirement of genome integration (4,5). Accomplishment of siRNA mediated specific gene silencing requires its effective delivery into target cells. (6–8). There are possibly thousands of materials that can be complexed with siRNA for its delivery into cells. For example, lipid based liposomal nanoparticles, PEI based complexes, calcium based complexes and AuNPs and other metallic NP based complexes (9–11). Out of these, choice must be made to select a material with effective complexing abilities with nucleic acids and must be biocompatible and less cytotoxic. AuNPs serve as a striking tool for nucleic acid delivery (12). Advantages of using AuNPs include their synthesis in required size and other metallic NP based complexes

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lize AuNPs to fabricate a complex of calcium and PEI in layer by layer assembly to deliver siRNA against HER2 into breast cancer MCF-7 cell line. We demonstrate the successful delivery of siRNA through AuNPs and calcium-based formulation assembly for knocking down the expression of HER2 gene.

Materials and Methods

Synthesis and characterization of AuNPs

Gold nanoparticles were synthesized by citrate reduction method (23). 20 mL of 1.0 mM hydrogen tetrachloroaurate (HAuCl₄) was boiled, 2 mL of 1% solution of trisodium citrate dihydrate (Na₃C₆H₅O₇·2H₂O) was added to boiling solution of HAuCl₄. After 10 minutes of boiling, citrate reduced gold ions and AuNPs were produced turning suspension in deep cherry red color. Particles were characterized by scanning electron microscopy (SEM) (Joel instruments) and dynamic light scattering (DLS) method via Nanosizer (Malvern instruments) for size, hydrodynamic diameter and surface charge.

siRNA and primer sequences

Sequence of HER2-siRNA and PCR primers used in this study are shown in table 1.

Layer by layer assembly of HER2-siRNA/AuNPs formulation

HER2 sense and antisense siRNA were re-suspended and annealed in annealing buffer, 0.1 mg/mL solution of 11-MUA, 0.5 mg/mL solution of PEI (25kDa, Branched), 2 M solution of CaCl₂, were made separately and filtered sterilized using 0.2 µM syringe filters. For the first layer of coating, 0.2 mg/mL NPs were obtained in 1.5 mL centrifuge tube in total volume of 500 µL. 11-MUA was added at 0.1 mg/mL, incubated for 30 minutes at room temperature and centrifuged at 16000 x g twice for 5 minutes to remove unbound 11-MUA and mixed in 100 µL, 10 mM NaCl solution. For the second layer, 2 M CaCl₂ was added to AuNPs from previous step and incubated for 30 minutes followed by centrifugation at 16000 x g twice for 5 minutes to remove unbound CaCl₂ and mixed in 100 µL, 10 mM NaCl solution. For the third layer, 2.1 µM HER2-siRNA was added to AuNPs from previous step and incubated for 30 minutes followed by centrifugation at 16000 x g twice for 5 minutes to remove unbound siRNA and mixed in 100 µL, 10 mM NaCl solution. For fourth and final layer of HER2-siRNA/AuNPs formulation, 1.0 mg/mL PEI was added to AuNPs from previous step and incubated for 30 minutes followed by centrifugation at 16000 x g twice for 5 minutes to remove unbound PEI and mixed in 100 µL, 10 mM NaCl solution (Fig. 1). After addition of each layer, the formulation was subjected to spectral scan via UV-VIS spectrophotometry to determine the peak plasmon shift. ζ potential was also determined after addition of each layer to confirm the charge reversal by each polymer.

siRNA release profile

After the preparation of NPs formulation, the release profile of siRNA was characterized. In order to determine the release profile, 1 mL of formulation was incubated in 3 mL of phosphate buffer saline (PBS, pH 7.3-7.4) at 37 °C at different time intervals for up to 144 hours. The samples were centrifuged at 13,000 x g for 30 min at 37°C and resulting pellets were re-suspended in 3 mL of PBS after each designated time interval. The supernatant from each centrifugation step was used to check for the presence of released siRNA by measuring absorbance at 260 nm via UV spectrophotometry. The amount in the supernatant was then subtracted from the initial amount loaded to determine the amount released at a given time point.

Cytotoxicity assay

MCF-7 cells were grown in Dulbecco’s Modified Eagle Media (DMEM), with 10% fetal bovine serum (FBS) and 1% penicillin/streptomycin, 0.1 mM L-glutamine, 1.0 mM sodium pyruvate with 2.4 g/L sodium bicarbonate in T25 cell culture flasks in a humidified incubator at 37°C. After sub culturing the cells were plated in 24 well plate at cell count of 2.5 10⁴ cells per well for transfection studies and incubated in humidified CO₂ incubator at 37°C. 100 µL of complex was mixed with serum free 150 µL medium. Mix was incubated for 5 minutes before adding to the cells after removing the older complete growth medium. After addition of complex in triplicates, plates were incubated in humidified incubator at 37°C for 48 hours. MTT cytotoxicity assay was performed according to instruction provided by manufacturer.

RNA isolation and cDNA synthesis

MCF-7 cells were separately transfected with AuNPs formulation. After 48 hours of transfection, cells were trypsinized, washed with phosphate buffer saline (PBS)
and collected in micro centrifuge tubes. Total RNA was isolated using TRIzol® Plus RNA Purification Kit from Thermo Fisher according to protocol. Purified RNA was stored at -80°C. cDNA was synthesized by Maxima First Strand cDNA Synthesis Kit (Thermo Fisher) according to the manufacturer protocol.

Gene expression profiling by quantitative real-time PCR

To determine the level of mRNA in transfected and untreated cells, cDNA was subjected to qPCR using 5x HOT FIREPol® EvaGreen® qPCR Mix Plus according to the manufacturer protocol with following thermal profile: initial activation for 12 min at 95°C, denaturation for 30 seconds at 95°C, annealing for 30 seconds at 52°C for HER2 and 57°C for ACTB respectively, elongation for 30 seconds at 72°C, thermal profile was run 40 times from denaturation to elongation. The green/FAM channel was selected for EvaGreen fluorescence detection. The Ct values data were subjected to 2⁻ΔΔCt analysis to find out fold change.

Results

Characterization of nanoparticles

The average hydrodynamic diameter of spherical AuNPs measured by DLS was 34.44 nm and the polydispersity index (PDI) was 0.117 ± 0.002 as shown in histogram (Fig. 2).

The surface charge or zeta (ζ) potential was -12 ± 1.75 mV. AuNPs were subjected to UV-VIS spectrophotometry to obtain spectral scan which showed maximum absorption peak at 521 ± 2 nm (Fig. 3), thus confirming the presence of gold in suspension. Finally, SEM images confirmed the presence of AuNPs with an average diameter of 21.8 ± 1.3 nm (Fig. 4). It is worth noting that NPs size varied when measured with nanosizer, this is because of the difference in the working principles of these instruments. Nanosizer measures the hydrodynamic diameter by light intensity scattered by an object/nanoparticle in light path and average size is calculated based upon solution dynamics of particles. On the other hand, SEM yields images of individual particles and it is considered more suitable for metallic nanoparticles (24).

Layer by layer assembly of HER2-siRNA/AuNP complex

AuNPs were coated with 11-MUA, Ca, siRNA and PEI successively. Addition of each charge bearing successive layer resulted in the shift of plasmon peak from 520 nm to 535 nm (Fig. 5) which indicated the addition of respected layer. Furthermore, the intensity of absorbance of bare AuNPs was reduced from 0.087 to 0.05 which showed that the AuNPs are being covered evenly by layers of charged polymers (Fig. 6). The surface charge analysis of each successive layer revealed that bare AuNPs had slight negative charge of -1 to -5 mV due to presence of negatively charged citrate ions. The addition of 11-MUA resulted in the even distribution of negative charge in the surface of AuNPs with -17 mV. Ca rendered the formulation with +19 mV charge while addition of negatively charged HER2-siRNA contributed to -20 mV and final layer of PEI provided +23 mV surface charge which would facilitate the entry of formulation in MCF-7 cells (Fig. 7).
siRNA release profile

Release profile of formulation was assessed by measuring the absorbance of released siRNA in supernatant at 260nm via UV-Vis spectrophotometry. siRNA release was assessed at different time intervals of 6, 12, 24, 48, 96, 120 and 144 hours. The results indicated that after 6 hours of incubation 60% siRNA was released from the formulation. After 12 hours of incubation 76% and after 24 hours 82% siRNA was released. This showed that the significant amount (more than 80%) of siRNA was released in early few hours. In the later hours the siRNA release was at much slower rate and 95% of siRNA was released up till 144 hours (Fig. 8). This quick liberation of siRNA might be attributed to the fact that Ca is a divalent ion and the electrostatic interactions between Ca ions and siRNA are relatively weaker (25,26).

Assessment of cellular uptake of formulation into cancer cells with EGFP plasmid

In order to assess the cellular uptake of formulation in MCF-7 cells, EGFP plasmid (Addgene plasmid #13031) was coated on AuNPs instead of HER2-siRNA just to confirm the entry of fabricated particles into the cells. EGFP plasmid was conjugated with lipofectamine2000 to serve as positive control for cellular uptake in MCF-7 cells and EGFP plasmid with no carrier was used to serve as negative control. After 48 hours of transfection with AuNP-EGFP, positive and negative control, MCF-7 cells were fixed using 4% paraformaldehyde and imaged with fluorescence microscope for the presence of fluorescence produced by green fluorescence protein (GFP) expressed by EGFP plasmid. Our results indicated the presence of fluorescence of GFP in both positive control (Fig. 9) and AuNP-EGFP formulation (Fig. 10). These results indicated that our formulation is capable of carrying nucleic acids into the cells.

Cytotoxicity assessment of formulation

MCF-7 cells were transfected with variable concentrations of NPs formulation and lipofectamine2000 conjugated with HER2-siRNA serving as positive control to assess the cytotoxicity. HER2-siRNA alone with no carrier served as negative control. The results showed decrease in the viability of cells at various concentrations. In lipofectamine2000 formulation cell viability decreased from 100% to 20% with concentration of 10 nM to 70 nM. Increase in the concentration of beyond 70 nM did not show any notable change. The half maximum inhibitory value (IC$_{50}$) was calculated to be 49.94 nM ± 0.1 (n=3). (Fig. 11a, 11b). The cells treated with siRNA conjugated with AuNPs in Ca formulation showed the dose dependent decrease in the cell viability. Viability assessment of AuNPs with HER2-siRNA treated cells showed that at 10nM concentra-
tion, 97.11% cells were viable which were decreased to 85% at the concentration of 20nM. An obvious change in decreased viability was found at 30nM concentration when 58% cells were found viable. Viability of cells was further decreased to 36.5% at concentration of 40nM. When concentration was increased to 50nM, viability was further decreased to 26.7% and it became 22.2% at the concentration of 60nM. Little bit decrease in viability was observed at the concentration of 70nM i.e. 20%. It could be seen from results that an obvious decrease has been found between concentrations of 20nM to 40nM when viability was decreased from 85% to 36.5%. Beyond 60nM observed change was negligible (Figure 12a, 12b). The IC<sub>50</sub> was 45.35nM ± 0.1 (n=3). These results indicated that the HER2-siRNA/AuNPs formulation has slightly higher cytotoxicity at relatively lower concentration as compared to positive control.

**Gene expression analysis**

The expression analysis on MCF-7 cell treated with 100 nM (2X of IC<sub>50</sub>) of HER2-siRNA/lipofectamine2000 (positive control) and 91 nM (2X of IC<sub>50</sub>) of HER2-siRNA/AuNPs was performed. Results of cells treated with positive control showed that 2^-ΔΔCt values for the treated and untreated cells were 0.152 and 1.006 respectively. This revealed a significant decrease (p<0.001) in the expression of HER2 gene when compared to untreated cells and fold decrease in the expression of HER2 was 6.61 times. (Fig. 13). On the other hand, MCF-7 cell treated with HER2-siRNA/AuNPs showed 2^-ΔΔCt values of 1.004 and 0.053 in treated and untreated cells respectively. This also led to the significant decrease (p<0.001) in the expression of HER2 by 18.94 times (Fig. 14).

**Discussion**

RNAi mediated gene silencing has appeared as a powerful strategy in handling cancer producing genes by complementary base-pairing mechanism (27). siRNA is a very important tool with substantial use in cancer therapy (28). Specifically, designed siRNA has the ability to modify the expression of cancer-causing
genes by homology-based pairing and post-transcriptional silencing. siRNA is a powerful tool to knockdown the oncogenes due to its specific and efficient silencing of target mRNAs (29). As previously reported, the efficiency and specificity of siRNA could be enhanced with the help of NPs as carriers (30), which have emerged as a striking tool for the delivery of biomolecules due to their easy synthesis, good biocompatibility, ready functionalization and monodispersity (31,32). This could be clearly observed in our results. The formulation successfully killed MCF-7 cells with an IC$_{50}$ of 45.35 nM while the IC$_{50}$ of siRNA conjugated with lipofectamine2000 was 49.94 nM. This indicates a higher degree of cell death when siRNA was conjugated in layer by layer assembly, probably due to the slow and steady release and greater half-life of siRNAs that resulted in suppression of MCF-7 cells proliferation. Furthermore, in terms of gene expression analysis, our results showed that with lipofectamine2000, the expression of HER2 was reduced by 6.1 times, while HER2-siRNA/AuNPs complex reduced the expression of HER2 by 18.94 times, indicating a greater efficiency of AuNPs in siRNA delivery to the cells. These results are consistent with previous finding which showed that AuNPs, due to inert character of metallic gold are non-toxic, form fine NPs with easy functionalization provide safer and efficient means of siRNA delivery (33). These results also confirm the efficiency of siRNA in knockdown of oncogene with AuNPs delivery system, indicating the therapeutic role of siRNA and NPs in cancer models (34,35). Furthermore, the previous studies showed that after introduction of the layer by layer technology for siRNA-AuNPs delivery into the cell by Elbakry et al. (2009). PEI was used as a gold standard for gene delivery but PEI is cytotoxic itself. (36,37). Due to additional cytotoxicity conferred by PEI, we used calcium in our formulation as AuNPs/MUA/Ca/siRNA/PEI. The application of calcium-based biomaterials in gene delivery causes calcium ion to form ionic complexes with the helical phosphates of DNA, and these complexes have easy transportability across the cell membrane via ion channel-mediated endocytosis. (38). Our results indicated that this formulation worked efficiently.

In conclusion, The present study shows calcium-based layer by layer assembly of siRNA-AuNPs silencing complex which successfully delivered HER2-siRNA into MCF-7 cells and efficient knockdown of HER2 gene. The RT-PCR based delta-delta Ct analysis in treated cells revealed a significant 18.94 times decrease ($p<0.001$) in the expression of HER2 gene standardized with ACTB housekeeping gene as compared to untreated cells, which makes this formulation a promising approach. Based on the results it is suggested that in future, the formulation should be tested in the xenographed mice models to evaluate its delivery and knockout potential in-vivo.

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Conflict of Interest
The Authors declare no conflict of interest for this publication.

References
Calcium based siRNA coating on gold nanoparticles.


