Potential inhibition of HIV-1 encapsidation by oligoribonucleotide–dendrimer nanoparticle complexes

Raveen Parboosing1,2
Louis Chonco1,2
Francisco Javier de la Mata3,4
Thavendran Govender5
Glenn EM Maguire5
Hendrik G Kruger5

1Department of Virology, University of KwaZulu-Natal, 2National Health Laboratory Service, Durban, South Africa; 3Organic and Inorganic Chemistry Department, University of Alcalá, Alcalá de Henares, 4Networking Research Center on Bioengineering, Biomaterials and Nanomedicine (CIBER-BBN), Madrid, Spain; 5Catalysis and Peptide Research Unit, University of KwaZulu-Natal, Durban, South Africa

Background: Encapsidation, the process during which the genomic RNA of HIV is packaged into viral particles, is an attractive target for antiviral therapy. This study explores a novel nanotechnology-based strategy to inhibit HIV encapsidation by an RNA decoy mechanism. The design of the 16-mer oligoribonucleotide (RNA) decoy is based on the sequence of stem loop 3 (SL3) of the HIV packaging signal (Ψ). Recognition of the packaging signal is essential to the encapsidation process. It is theorized that the decoy RNA, by mimicking the packaging signal, will disrupt HIV packaging if efficiently delivered into lymphocytes by complexation with a carbosilane dendrimer. The aim of the study is to measure the uptake, toxicity, and antiviral activity of the dendrimer–RNA nanocomplex.

Materials and methods: A dendriplex was formed between cationic carbosilane dendrimers and the RNA decoy. Uptake of the fluorescein-labeled RNA into MT4 lymphocytes was determined by flow cytometry and confocal microscopy. The cytoprotective effect (50% effective concentration [EC50]) and the effect on HIV replication were determined in vitro by the methylthiazolyltetrazolium bromide (MTT) assay and viral load measurements, respectively.

Results: Flow cytometry and confocal imaging demonstrated efficient transfection of lymphocytes. The dendriplex containing the Ψ decoy showed some activity (EC50 = 3.20 μM, selectivity index = 8.4). However, there was no significant suppression of HIV viral load.

Conclusion: Oligoribonucleotide decoys containing SL3 of the packaging sequence are efficiently delivered into lymphocytes by carbosilane dendrimers where they exhibit a modest cytoprotective effect against HIV infection.

Keywords: packaging signal, dendrimers, transfection, antiretroviral, HIV packaging

Introduction

Encapsidation, the process by which the dimeric RNA of HIV is preferentially packaged by the Gag polyprotein into nascent virions, is an essential step in HIV-1 replication.1-4 This process involves interaction between the gag nucleocapsid protein p7 (NCp7) and a region of the HIV genome known as the packaging signal (Ψ). This interaction is an attractive target for antiviral therapy because the secondary structures of Ψ and nucleocapsid are highly conserved among retroviruses5-9 and mutations in NCp7 result in totally defective virions.10-12 This suggests that drug resistance to “packaging inhibitors” is less likely to develop. Furthermore, RNA export (ie, packaging) is a virus-specific and not a cellular process so that drugs targeting this step are likely to have a high therapeutic index.1

Studies exploring the potential of targeting the packaging process have shown promising results. For example, vectors that express Ψ-containing transcripts,
or constructs that contain the packaging signal, interfere with packaging and successfully inhibit HIV replication in vitro. In another study, RNA ligands (“aptamers”) with Ψ-like sequences bound to NCp7 with high affinity and abrogated packaging (where the ligand, rather than the genome, is packaged, resulting in defective virions). Antisense RNA that targets the packaging signal inhibits retroviral replication.

Four “stem-loops”, SL1–4, constitute the secondary structure of Ψ; of these, SL3 has been identified by mutational, biochemical, SL-NCp7 affinity, ligand substitution and nuclear magnetic resonance (NMR) experiments to play a particularly important role in viral RNA packaging. The aim of this study is to design and optimize a potential oligoribonucleotide antiviral therapeutic that will inhibit packaging by several possible mechanisms such as, 1) competitive inhibition: the oligoribonucleotide will compete with SL3 (and other SLs) for binding sites on NCp7; 2) decoy encapsidation: packaging of the oligoribonucleotide decoy, rather than the HIV genome, will reduce the pool of available NCp7 for Ψ binding and packaging; 3) defective interfering particles; and 4) interference with the role of NCp7 (especially as a “nucleic acid chaperone”) in other steps of replication, such as initiation of reverse transcription, strand transfer, integration, gag multimerization, RNA dimerization and viral assembly.

Lymphocytes, in particular, and suspension cell lines, in general, are, however, notoriously difficult to transfect. Nonetheless, there have been reports of successful transfection of lymphocytes using carbosilane dendrimers. This prompted us to investigate the potential of a carbosilane dendrimer to deliver the SL3 oligoribonucleotide decoy into CD4+ lymphocytes, thereby inhibiting the replication of HIV.

Materials and methods

Reagents

The following reagents were obtained through the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH: MT4 from Dr Douglas Richman and HIV PMC/H9 from Dr Robert Gallo. The RNase-free, high-performance liquid chromatography (HPLC)-purified RNA construct was synthesized by Dharmacon (Pittsburgh, PA, USA). BLOCK-i™ Fluorescent Oligo (used as control RNA) was obtained from Thermo Fisher Scientific (Waltham, MA, USA). Nuclease-free water was obtained from Qiagen (Hilden, Germany). Heat-inactivated fetal bovine serum (FBS) was obtained from BioChrom/Merck Millipore (Berlin, Germany). OptiMEM™ and Gibco® fetal bovine serum (FBS) was obtained from Life Technologies (Carlsbad, CA, USA). Phosphate-buffered saline (PBS) without Ca/Mg was obtained from Lonza (Basel, Switzerland). All other reagents were obtained from Sigma unless otherwise indicated.

Design of the RNA decoy

An oligoribonucleotide decoy containing a 16-mer sequence of SL3 (ACUAGCGAGGCUCUAGA) was synthesized for use in this study (Figure 1). For uptake studies, the RNA was modified to include a fluorescein label at the 3’ end. The design of the RNA decoy was based on, 1) the structure of Ψ, and its molecular interaction with NCp7, including NMR spectroscopy elucidation of its three-dimensional conformation, evidence from insertional mutagenesis, ligand binding, and biochemical experiments regarding the factors (eg, the nucleotide sequence and content) that determine the affinity between Ψ and NCp7, 3) previous in vitro gene therapy experiments (using either aptamers or vectors) in which Ψ-like nucleotide sequences have successfully abrogated packaging, and 4) previous studies using archetypal SL3 ligands, Systematic Evolution of Ligands by Exponential Enrichment (SELEX) ligands and synthetic RNA stem loops.

RNA preparation and handling

The custom-made RNA (16-mer sequence of SL3) was deprotected as recommended by the manufacturer, dried using a centrifugal evaporator (mIVac DNA Sample Concentrator; Genevac, Stone Ridge, NY, USA), stored at −80°C as a pellet and resuspended in RNase-free deionized water just prior to use. RNase-free consumables were used, and precautions were taken to keep surfaces free from RNase (RNaseZap® wipes from Thermo Fisher Scientific). The RNA was quantified by measuring absorbance at 260 nm (adjusted for the effect of fluorescein) (BioSpec Nano; Shimadzu Corporation, Kyoto, Japan).

Dendrimer synthesis

A third-generation cationic carbosilane dendrimer was used for this study. This dendrimer contains 24 positive charges on
its skeleton and terminal –OH groups that could participate in hydrogen bonds with biomolecules (Figure 2). The synthesis of this dendrimer has been previously reported and involves the initial synthesis of a carbosilane dendrimer decorated with terminal dimethyl amino groups and posterior quaternization with 2-iodoethanol.32

**Dendriplex formation and optimization**

Dendriplexes were formed by mixing equal volumes of dendrimer and RNA (dissolved in RNase-free, deionized water), at varying molar ratios, and incubating for 15 minutes at room temperature, away from light. Freshly prepared dendriplexes (<1 hour old) were used to treat the cells. The ratio of dendrimer:RNA (± charge ratio) and the dose of the dendriplex were optimized in a series of experiments, based on the degree of complexation (gel retardation assay), uptake (flow cytometry and confocal microscopy), cytotoxic vs cytoprotective effect (methylthiazolyldiphenyl-tetrazolium bromide [MTT] assay) and effect on viral replication (viral load). The dendriplexes used in the uptake (flow cytometry and confocal microscopy) experiments were prepared by mixing 10 µM RNA with the following serially diluted concentrations of dendrimer (bearing in mind that the dendrimer has 24 positive charges and the RNA 16 negative charges per molecule, ie, P/N=3:2): dendriplex (8:1): 54 µM; dendriplex (4:1): 27 µM; dendriplex (2:1): 13.5 µM and dendriplex (1:1): 6.75 µM. In the MTT assay, serial dilutions of the dendrimer or dendriplex were used, beginning with the highest concentration of 75 µM. In experiments where the control RNA was used, the concentration of dendrimer and ± charge ratio was equivalent to that of the Ψ RNA.

**Gel retardation assay**

Complexation was studied in 2% agarose gel electrophoresis, using fluorescently labeled RNA and loading buffer.33

**Cells and virus**

MT4 lymphocytes were grown in RPMI 1640 medium supplemented with 10% heat-inactivated FBS at 37°C in 5% CO₂. Cells were used at the exponential phase of growth and with viability at least 95%. The Countess™ automated Cell Counter (Thermo Fisher Scientific, Waltham, MA, USA) was used to determine cell counts and viability.

Stock virus was prepared by harvesting supernatant of HIV-infected MT4 lymphocytes at day 5 post-infection. The virus was titrated by conventional methods and stored at −80°C until use.53 All work involving HIV culture was performed in the appropriate biosafety conditions.54,55

![Figure 2 Third-generation cationic carbosilane dendrimer used in this study.](image-url)
Flow cytometry
MT4 lymphocytes (150,000 cells/well) were suspended in wells of a microtiter plate with 200 µL OptiMEM™. The cells were then incubated for 4 hours with 25 µL of untreated control (OptiMEM™ only), RNA control (RNA only), dendrimer only (54 µM), 10% dimethyl sulfoxide (DMSO; dead cell control) and dendriplexes at various molar ratios. The cells were harvested, washed first with PBS, then with acid–glycine buffer for 30 seconds and finally again with PBS. The acid–glycine buffer (for the removal of dendriplexes bound to the surface of the cell membranes) consisted of 0.2 M glycine in PBS (adjusted to pH 3.0 by addition of hydrochloric acid).

The cells were then fixed with paraformaldehyde (at 4°C for 15 minutes) and then finally washed and resuspended in PBS. The cells were pelleted by gentle centrifugation after each wash step. The cells were stored at 4°C until analysis by flow cytometry (within 24 hours). The experiment was repeated without the acid–glycine wash to estimate the proportion of surface-bound vs intracellular RNA.

Uptake and viability were determined on BD FACS-Canto™ II (BD Biosciences, San Jose, CA, USA) instrument. To determine viability, 2 µL of Zombie Aqua™ dye (Biolegend, San Diego, CA, USA) was added to cell suspensions prior to fixation, as per manufacturer’s instructions. Uptake of RNA was quantified as the percentage of cells that took up fluorescein.

Confocal microscopy
Cells were treated, harvested and washed as for flow cytometry and then added to poly-L-lysine-coated microscope slides. The slides were fixed with paraformaldehyde and incubated for 10 minutes with Fluoroshield™ with 4′,6-diamidino-2-phenylindole (DAPI) stain and then viewed and photographed under confocal microscope, ZEISS Confocal LSM 710 (Zeiss, Oberkochen, Germany).

MTT assay
The MTT assay to determine cytotoxicity (50% inhibitory concentration [IC₅₀]), cytoprotective (antiviral) effect (50% effective concentration [EC₅₀]) and selectivity index (SI) was performed in triplicate as previously described. Briefly, MT4 lymphocytes were seeded in wells of a microtiter plate at 6x10⁵ cells/mL and treated with serial dilutions of azidothymidine (AZT), dendrimer alone, dendriplex with control RNA or dendriplex with Ψ RNA. A total of 50 µL of RPMI 1640 medium was added to each well in one half of the plate (mock infection), while 50 µL of HIV IIIb at 300 tissue culture infective dose (TCID₅₀) was added to the other half (HIV infection). The plate was incubated for 5 days at 37°C in 5% CO₂. At the end of the incubation period, the MTT assay was performed to assess toxicity in mock (uninfected) cells and cytoprotective effect in HIV-infected cells. The optical density readings were plotted on a graph, and the IC₅₀ and EC₅₀ were determined by linear extrapolation.

Viral load assay
MT4 cells were incubated for 3 hours with HIVIIIb at multiplicity of infection (MOI) 0.01, washed thrice with PBS to remove the initial inoculum and then plated at 3x10⁴ cells/mL (200 µL per well with RPMI 1640 medium and 10% FBS). Wells were treated in triplicate with 25 µL of AZT, dendrimer alone, dendriplex with control RNA or dendriplex with Ψ RNA. The concentration of the dendrimer was kept constant. A total of 25 µL of RPMI 1640 medium was added to untreated control wells. After 3 or 5 days of incubation, 100 µL was aspirated from each well, inactivated by exposure to acidified Triton X (in isopropanol) for 1 hour, washed twice (by centrifugation and resuspension in PBS) and finally resuspended in 3 mL PBS for viral load quantification. The dilution factor (10⁴) introduced because of the aforementioned procedures was taken into account when calculating the final viral load. Viral load was quantified by the automated Roche Cobas® 6800/8800 System (Roche Diagnostics, Mannheim, Germany) as per the manufacturer’s instructions. This is a fully automated polymerase chain reaction (PCR)-based system with a linear range of 20–10,000,000 copies/mL.

Statistical analysis
Unpaired Student’s t-test was used to calculate P-values. A P-value of <0.05 was regarded as significant.

Results
The oligoribonucleotide molecule formed complexes with the carbosilane dendrimer
Successful complexation was demonstrated on gel electrophoresis when the molar ratio of dendrimer:RNA exceeded 2 (corresponding to a ± charge ratio of >3:1) (Figure 1). Faint bands appear at lower ratios, suggesting partial complexation (Figure 3).
The carbosilane dendrimer facilitated efficient transfection of the oligoribonucleotide decoy into MT4 lymphocytes

The gating for fluorescein uptake was adjusted so that the untreated control showed <1% fluorescence. The dendriplex with ± charge ratio of 4:1 and 8:1 demonstrated efficient transfection of MT4 lymphocytes on flow cytometry (90.1% and 98.5%, respectively). Dendriplexes with lower charge ratios (2:1 and 1:1) achieved lower transfection (15.7% and 1.9%, respectively) (Figure 4). By comparison, in other studies, transfection efficiencies of 36% and 90% were reported with the use of a second-generation carbosilane dendrimer in primary T lymphocytes\textsuperscript{33} and SupT1 lymphocytes,\textsuperscript{34} respectively. The RNA alone was taken up by 9.3% of cells.

Removal of surface-bound dendriplexes by an acid–glycine wash suggests that the uptake results represent complexes located within the cell (rather than on the surface of the cell membrane). Omission of the acid–glycine wash resulted in only slight differences in the fluorescence reading (97.6% and 92.6% vs 90.1% and 98.5% for the 4:1 and 8:1 dendriplexes, respectively), suggesting that most of the fluorescence represents intracellular RNA.

Dead cells were identified by staining with Zombie Aqua\textsuperscript{TM} dye. Debris was gated out by its characteristic appearance on side scattered light (SSC)-A and forward scattered light (FSC)-A. The gating was applied so that the Zombie Aqua\textsuperscript{TM} stain correctly determined the viability of untreated cells (∼95%) (compared to Trypan Blue staining). Toxicity, based on this gating strategy, was determined to be as follows: RNA only (10 µM): 9.5%; dendrimer only (54 µM): 76.5%; dendriplex (8:1): 20.7%; dendriplex (4:1): 10.4%; dendriplex (2:1): 6.4% and dendriplex (1:1): 3.2%.

Uptake was confirmed by confocal microscopy, which furthermore showed the cytoplasmic location of the fluorescent RNA (Figure 5). The dendrimer with charge ratio 2:1 did not show any uptake by confocal microscopy.
Cytotoxic and cytoprotective effect of dendriplex and controls

The dendriplex containing the Ψ RNA exhibited a cytoprotective effect and selectivity (IC$_{50}$ = 26.8 μM, EC$_{50}$ = 3.20 μM and SI = 8.4), while the RNA alone and the dendriplex containing control RNA had no cytoprotective effect (Figure 6; Table 1). The dendriplex with the Ψ RNA restored infected cells to ~60% of the viability of uninfected, untreated cells (Figure 6).

MT4 lymphocytes exposed to the dendriplex retained >80% viability at a dendrimer concentration of 15 μM (equivalent to 138 μg/mL). By comparison, peripheral blood mononuclear cells and SupT1 cells exposed to a second-generation carbosilane dendrimer in a study by Weber et al.$^{26}$ reached the toxicity limit of 80% at 24 μg/mL, while in another study, two second-generation dendrimers reached the toxicity limit for macrophages at 5 and 15 μM, respectively.$^{33}$

Viral load assay

HIV-infected lymphocytes treated with the dendriplex did not demonstrate significant reduction in viral load following 3 days of incubation. There was no difference in viral load between the dendriplex containing the Ψ decay vs control RNA or dendriplex alone without RNA (data not shown). The same result was obtained when the incubation period of the experiment was increased to 5 days.

Discussion

This study showed efficient uptake of a dendrimer–oligoribonucleotide nanocomplex into MT4 lymphocytes with evidence of a cytoprotective effect (EC$_{50}$ = 3.20 μM, SI = 8.4) against HIV. However, the dendriplex did not demonstrate significant suppression of HIV viral load. There may be several reasons why the oligoribonucleotide decoy did not have a more profound antiviral and cytoprotective effect. First, interaction of the nucleocapsid protein (NCp7) with the encapsidation signal possibly depends on secondary intermolecular and intramolecular structural motifs rather than a consensus sequence.$^{1,2,4}$ It is not known how well a decoy based on a linear sequence will simulate these motifs. Further studies are required whereby the sequence

![Table 1](https://www.dovepress.com/)

**Table 1** Toxicity (IC$_{50}$) and cytoprotective effect (EC$_{50}$) of Ψ RNA dendriplex and various controls

<table>
<thead>
<tr>
<th>Treatment</th>
<th>IC$_{50}$ (μM)</th>
<th>EC$_{50}$ (μM)</th>
<th>SI (IC$<em>{50}$/EC$</em>{50}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AZT (antiviral control)</td>
<td>13.26</td>
<td>0.0023</td>
<td>5,766</td>
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<tr>
<td>Dendriplex</td>
<td>26.8</td>
<td>3.20</td>
<td>8.4</td>
</tr>
<tr>
<td>(Ψ RNA + dendrimer)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RNA alone</td>
<td>Nontoxic*</td>
<td>No effect*</td>
<td>Not calculable*</td>
</tr>
<tr>
<td>Dendrimer alone</td>
<td>26.4</td>
<td>No effect*</td>
<td>Not calculable*</td>
</tr>
<tr>
<td>Dendriplex</td>
<td>21.7</td>
<td>No effect*</td>
<td>Not calculable*</td>
</tr>
<tr>
<td>(control RNA + dendrimer)</td>
<td></td>
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</tbody>
</table>

**Notes:** *In the dosage range tested.

**Abbreviations:** IC$_{50}$, 50% inhibitory concentration; EC$_{50}$, 50% effective concentration; SI, selectivity index.
and number of nucleotides that constitute the decay are varied to determine the effect of decay nucleotide structure on packaging efficiency. Second, there is probably redundancy in NCp7-Ψ interaction,24,25,59 ie, more than one stem loop may be able to initiate packaging. To explore this, all four stem loops, singly and in combination, should be evaluated for their anti-packaging effect. Finally, stoichiometry between NCp7 and Ψ may make inhibition challenging. There are ≥1,500 gag molecules per dimerized viral genome.56,16,59,60 The number of Ψ decoys required to effectively interfere with packaging is not known. Although the dendriplex achieved excellent transfection efficiency, this is not a direct measure of the intracellular Ψ RNA dose (which is limited by the toxicity of the carrier molecule and ± charge ratios). Furthermore, it is not known whether the RNA is able to exert its activity while bound to the dendrimer, and if not, what proportion is released from the dendriplex once it reaches the cytoplasm.

The cytoprotective effect of the dendriplex was not corroborated by the viral load experiment. This paradoxical outcome could be an artifact of the experimental setup. Further mechanism of action studies will be required to find a suitable explanation. Furthermore, it will be of interest to determine if this finding is replicated with other dendrimers and alternative transfection methods.

We did not study the mechanism of uptake of the dendriplex, which is known to be cell type dependent.61 Furthermore, we did not assess the rate of uptake, which may be influenced by the cationic nature of the dendrimer.62 These are avenues for further research, particularly in lymphocytes that are naturally resistant to transfection.

Conclusion

While this study showed evidence of the cytoprotective effect of a Ψ RNA decay dendriplex, potent inhibition of encapsidation by this strategy probably requires several issues to be addressed, including optimized design and intracellular delivery of the packaging signal decay. More importantly, this study provides further evidence that cationic dendrimers may be an efficient non-viral alternative for the delivery of anti-HIV RNA therapeutics into hard-to-transfect lymphocytes.

Acknowledgments

Open access publication of this article has been made possible through support from the Victor Daitz Information Gateway, an initiative of the Victor Daitz Foundation and the University of KwaZulu-Natal. Logan Reddy and staff in the Viral PCR laboratory at the National Health Laboratory Service in Durban for HIV viral loads, Kogi Moodley/Prof Daniels at the University of KwaZulu-Natal (UKZN) for use of flow cytometry equipment, and Lorika Beukes at UKZN for use of the confocal microscope, are acknowledged. University of Alcalá acknowledges financial support from CTQ2014-54004-P (MINECO) and Consortium NANO-DENDMED ref S2011/BMD-2351 (CM). CIBER-BBN is an initiative funded by the VI National R&D&I Plan 2008–2011, Iniciativa Ingenio 2010, Consolider Program, CIBER Actions and financed by the Instituto de Salud Carlos III with assistance from the European Regional Development Fund. Dr Parboosing received an Academic Fellowship award from the Discovery Foundation of South Africa and funding from the National Research Foundation Thuthuka Program.

Disclosure

The authors report no conflicts of interest in this work.

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