Novel Disulfide-Bridged Bioresponsive Antisense Oligonucleotide Induces Efficient Splice Modulation in Muscle Myotubes in Vitro

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ABSTRACT: Splice-modulating antisense therapy has shown tremendous potential in therapeutic development in recent years with four FDA-approved antisense drugs since 2016. However, an efficient and nontoxic antisense oligonucleotide (AO) delivery system still remains as a major obstacle in nucleic acid therapeutics field. Vitamin-E (α-tocopherol) is an essential dietary requirement for human body. This fat-soluble compound is one of the most important antioxidants which involves in numerous biological pathways. In this study, for the first time, we explored the scope of using α-tocopherol-conjugated bioresponsive AOs to induce splice modulation in mouse muscle myotubes in vitro. Our results showed that the bioresponsive construct efficiently internalized into the cell nucleus and induced exon 23 skipping in mdx mouse myotubes. Based on our exciting new results, we firmly believe that our findings could potentially benefit toward establishing a delivery approach to advance the field of splice-modulating AO therapy.

1. INTRODUCTION

Synthetic antisense oligonucleotide (AO) has attracted extensive interest in therapeutic development for various genetic diseases after demonstrating successful inhibition of Rous sarcoma virus 35S RNA translation in 1978.1 Since then, eight AO drugs have been approved for clinical use including Vitrivene, Kynamro, Tegsedi, and Waylivra utilizing RNase-H-dependent pathway and Exondys 51, Spinraza, Milasen, and Vitravene using the splice modulation mechanism.2-6 In order to achieve high therapeutic efficacy, AO drug molecules need to be stable under nuclease conditions while possessing high binding specificity and affinity to the RNA target. In addition, tissue distribution and cellular uptake properties of AOs also play a pivotal role in the overall treatment outcome. However, despite the dramatic progress in recent years, establishing an effective AO delivery system still remains as one of the major challenges. Various delivery approaches have been studied for AOs over the last few years, and a comprehensive review of this topic can be found elsewhere.7

Conjugation of biomolecules to AOs has attracted significant interest in therapeutic development because of the simplicity in the synthesis and pharmacokinetic studies. Polyethylene glycol was one of the first molecules utilized in the delivery of an aptamer drug Macugen.8,9 In addition, the potential of N-acetylgalactosamine (GalNAc)10 has been studied for liver-specific uptake of oligonucleotides, and based on this development, an siRNA drug called Patisiran11 has been approved by the US FDA that offers a great promise for liver-specific delivery of AOs. Furthermore, the potential of various cell-penetrating peptides has been studied extensively particularly to enhance the delivery of phosphorodiamidate morpholino oligo-based AO drug molecules.12

Recently, vitamin-E, a group of eight naturally occurring fat-soluble compounds, has emerged as a potential approach for oligonucleotide delivery.13,14 Vitamin-E, a widely used component in various cosmetic and dietary supplement products, has eight isoforms in which α-tocopherol (α-TOC, Figure 1) has been identified as the most abundant form in human body.15-16 α-TOC is believed to be beneficial for human health; however, there is no clear evidence on the positive effects as a treatment modality for disease prevention or therapy.16 Yokota and colleagues demonstrated the use of α-TOC for the delivery of siRNA and chimeric oligonucleotides α-TOC-conjugated heteroduplex oligonucleotide for the modulation of blood–brain barrier in mice.17-19 Another study from Kuwahara et al. also showed the efficacy of an α-TOC-conjugated heteroduplex oligonucleotide for the modulation of blood–brain barrier in mice.20 Inspired by these studies, we envisaged that the α-TOC-conjugated AO could also be applicable in the splice modulation scenario. However, unlike previous reports in which the AOs bind to the mature mRNA targets in the cell
cytoplasm, splice-modulating AOs need to be delivered into the cell nucleus where it binds to the pre-mRNA and interfere with splicing events. Toward addressing this challenge, we established a novel construct by introducing a disulfide bridge that links between the AO and α-TOC (α-TOC-S-S AO) (Figure 1) and evaluated its efficiency to induce exon 23 skipping in a dystrophin gene (Dmd) transcript model system by utilizing mdx mouse myotubes in vitro. The designed AO construct is bio-reducible inside the cell, which can promote and enhance the ability of the AO to efficiently internalize into the cell nucleus and induce exon skipping.

2. RESULTS AND DISCUSSION

To demonstrate this rationale systematically, we first assessed the effects of a non-S-S-containing AO construct to induce exon 23 skipping in vitro. Accordingly, a 20-mer α-TOC-conjugated 2′-O-methyl (2′-OMe, Figure 1) AO construct was designed and synthesized on a phosphorothioate (PS) backbone (α-TOC-AO, Table 1). The AO sequence used in this study was based on a previously reported best performing AO in inducing exon 23 skipping in mdx mouse myotubes.24 A non-conjugated 2′-OMePS was also synthesized as a positive control. We then evaluated the potential to induce exon 23 skipping by directly incubating the AO candidates with H-2K²-taSAS mdx (H2K mdx)25 mouse myotubes at four different concentrations (100, 200, 400, and 800 nM). In parallel, the 20-mer 2′-OMePS control AO was transfected with Lipofectin reagent at 100 nM. After 24 h, the cells were collected, and the total RNA was extracted, followed by performing nested reverse transcription polymerase chain reaction (RT-PCR) to amplify the products as previously described.21

As expected, the positive control yielded an efficient (62.3%) exon 23 skipping product of 688 bp (Figure S1; Supporting Information), in line with our previous observations.24−26 On the other hand, the α-TOC AO failed to induce any exon skipping after 24 h (Figure S1). Not surprisingly, the 2′-OMePS control AO without any lipids (naked form) was also not efficient after 24 h of incubation. Therefore, we increased the incubation time to enhance the uptake and repeated the experiment with prolonged incubation times of 48 and 72 h using the α-TOC AO.

Notably, at 48 h time point, although the 100 nM concentration failed to induce any exon skipping, other tested concentrations yielded the expected exon-23-skipped product in a dose-dependent manner ranging from 13.3% at 200 nM to 22.6% at 800 nM (Figure 2A,G). Similar to the 48 h time point, the exon 23 skipping products were observed at 72 h time point with all concentrations except 100 nM, and the skipping efficiency was similar between 200 and 400 nM (10.3 and 10.1%, respectively) and increased to 22.4% at 800 nM (Figure 2B,H). The decrease in the yield of exon 23 skipping compared with the 48 h time point could possibly be due to the unfavorable dual exon 22−23 skipping (542 bp product; 1.8−12.2% from 800 to 200 nM; Figure 2B,H) upon prolonged incubation time. It is also encouraging that the potency of the AO increased proportionally to the time of incubation. However, the efficiency of exon 23 skipping (688 bp product) was very low in comparison with the positive control 2′-OMePS AO transfected using Lipofectin. Simultaneously, an independent control experiment was also performed by incubating the Lipofectin transfection reagent-free (naked transfection) 2′-OMePS AO construct with the cells for 48 and 72 h. In this case, there was no exon skipping observed at 48 h (Figure 2E,K), while 72 h of incubation yielded exon 23 skipping in very low yields of 8.8 and 5.5% at 400 and 800 nM, respectively (Figure 2F,L), suggesting that a small amount of unconjugated AO could still be internalized and localized in the nucleus.

In line with our recent study (unpublished) using two different fatty-acid-conjugated AOs which failed to induce any exon 23 skipping, we speculated that α-TOC conjugation might also be limiting the nucleus localization of AOs after entering the cytoplasm. To investigate this hypothesis, we then explored the scope of a bioresponsive AO construct that can be cleaved in the cytoplasm by different mechanisms involving pH, glutathione level, or enzymatic reaction.28 For this purpose, incorporation of a reducible disulfide linkage (−S−S− linkage, Figure 1) could be a promising approach to cleave the AO from the α-TOC moiety. Higher concentration of glutathione (γ-glutamyl-cysteinyl-glycine; GSH) inside the cell compared to the extracellular matrix enables the cleavage of the disulfide bond by a redox reaction.29 Various studies, mainly in the field of cancer therapeutics, have utilized this biological mechanism for drug delivery. An update on this topic has been reviewed and published elsewhere.30 Notably, there were no previous reports on conjugating splice-
modulating AOs with a disulfide linkage. Therefore, in this study, we envisioned the scope of a novel bioresponsive AO construct (α-TOC-S-S-AO; Figure 1; Table 1) by linking α-TOC (the delivery agent) and the AO via a disulfide linkage (to facilitate AO release in the cytosol).

To validate this rationale, we incubated α-TOC-S-S-AO (Table 1) with the H2K mdx myotubes at 100, 200, 400, and 800 nM for 48 and 72 h. Overall, the bioresponsive α-TOC-S-S-AO induced far more efficient exon 23 skipping compared with the α-TOC-AO and lipid-free 2′-OMePS controls at both time points in line with our hypothesis. At 48 h, although 100 and 200 nM of α-TOC-S-S-AO failed to induce any skipping, higher concentrations yielded the skipped products in a dose-dependent manner, with 18% at 400 nM and 36.4% at 800 nM (Figure 2C,I). A dual exon 22−23 skipping product was also observed at 800 nM, which accounted for 18.3% of the total band intensity. Remarkably, 72 h of incubation yielded very efficient exon 23 skipping at 400 nM (46.7%), with a minimal amount of dual skipping product at 3.6% at 800 nM (Figure 2D,J). A dual exon 22−23 skipping product was also observed at 800 nM, which accounted for 18.3% of the total band intensity. Remarkably, 72 h of incubation yielded very efficient exon 23 skipping at 400 nM (46.7%), with a minimal amount of dual skipping product at 3.6% at 800 nM (Figure 2D,J). The results also showed that the exon-23-skipped product was slightly dropped to 40.6% at 800 nM, mainly because of the increase of the dual skipped product (26.4%). Not surprisingly, 200 nM of the AO only induced a slight exon 23 skipping (13.4%), while 100 nM failed to yield any skipped product. Nonetheless, the data were truly encouraging as it supported our hypothesis in designing the bioresponsive α-TOC-S-S-AO construct in which the disulfide linkage possibly played an important role in the nucleus localization of AOs.

In addition, to further demonstrate the validity of our results, we synthesized two fluorescent constructs α-TOC-AO-FAM and α-TOC-S-S-AO-FAM in which we labeled the AOs with a fluorescent dye (6-carboxyfluorescein) at the 3′-end of the sequence (3′-FAM; Table 1). The purpose of this experiment was to visualize the localization of the AO part inside the cell nucleus after the disulfide cleavage. Toward this, we incubated the AOs with the cells at 400 and 800 nM concentrations, and the images were taken at 72 h time point. Remarkably, our results indicated that both AOs were able to internalize into various myotube cell nuclei at both concentrations (Figure 3).
Quantification of the average fluorescence intensity in the nucleus (using ImageJ software) showed values of 72.1 and 56.1 for α-TOC-S-S-AO-FAM at 400 and 800 nM, respectively, in comparison to 20.5 and 40.3 of α-TOC-AO-FAM at 400 and 800 nM, respectively. Not surprisingly, this observation is in line with the exon skipping data.

3. CONCLUSIONS

In summary, our study has demonstrated the design and synthesis of a novel bioresponsive AO construct by conjugating α-TOC with a splice-modulating AO via a disulfide linkage (α-TOC-S-S-AO). The introduction of a bioreducible disulfide linkage certainly enhanced the uptake of the AO into the cell nucleus and therefore improved the exon 23 skipping potency in H2K mdx mouse myotubes. Based on our encouraging results, we firmly believe that these findings could certainly help toward establishing an efficient bioresponsive AO construct for inducing splice modulation.

4. MATERIALS AND METHODS

4.1. Design and Synthesis of AOs. All 2′-OMe (A, C, G, and U) phosphoramidites, DMT-6-FAM phosphoramidite, and α-TOC (vitamin E) phosphoramidite were purchased from ChemGenes (USA). S′-Thiol modifier C-6 disulfide modifier CED phosphoramidite was purchased from NOVASYNTH Technologies (I) LTD (India). Glen UnySupportTM (Universal solid support) was purchased from Glen Research (USA).

Modified PS oligonucleotides were synthesized as DMT-off in 1 μmol scale on a Glen UnySupport using the AKTA oligopilot system. Standard procedures were used for the coupling of commercial 2′-OMe phosphoramidites, whereas α-TOC (vitamin E) phosphoramidite, DMT-6-FAM phosphoramidite, and S′-thiol modifier C-6 disulfide modifier CED phosphoramidite were coupled with S′-(benzylthio)-1H-tetrazole (0.3 M) in CH3CN as an activator and an extended coupling time (20 min), the coupling efficiency was higher than 95% in all cases. Oligonucleotides were deprotected and cleaved from the solid support with 32% aq NH3 (1 mL) and left at 55 °C for 20 h. After deprotection, the oligonucleotides were desalted using an Illustra NAPTM-10 column (GE Healthcare). The purity of the final oligonucleotides was checked by ion-exchange chromatography using an HPLC system from Shimadzu on DNAPac PA200, a 250 × 4 mm analytical column. Buffers: [buffer A: MQ H2O; buffer B: 1 M NaClO4; and buffer C: 25 mM Tris-Cl, pH 8.0], flow rate: 1 mL min⁻¹.

4.2. Cell Culture and Transfection. H-2Kα-tsAS8 (H2K mdx) mouse myoblasts (provided by Prof. Sue Fletcher and Prof. Steve Wilton’s laboratory, Murdoch University, Australia) were cultured as described previously. Briefly, at 60–80% confluency, primary mdx myoblast cultures were treated with trypsin (Life Technologies) and seeded at a density of 2 × 10⁴ cells/well into 24-well plates. The plate was pretreated with 50 μg/mL of poly-o-lysine (Sigma) and 100 μg/mL of Matrigel (Corning). Cultures were induced to differentiate into myotubes in Dulbecco’s modified Eagle’s medium containing 5% horse serum by incubation at 37 °C in 5% CO₂ for 48 h. AOs were incubated directly with the cells, except the positive control which was complexed with Lipofectin (Life Technologies) at a ratio of 2:1 (Lipofectin/AO) and used in a final transfection volume of 500 μL/well in a 24-well plate as per the manufacturer’s instructions, except that the solution was not removed after 3 h.

4.3. RNA Extraction and RT-PCR. RNA was extracted from cells using the ISOLATE II RNA Mini Kit (Bioline) as per the manufacturer’s instructions. The dystrophin transcripts were then analyzed by nested RT-PCR across exons 20–26 as described previously. PCR products were separated on 2% agarose gels in Tris–acetate–EDTA buffer, and the images were captured on a Fusion Fx gel documentation system (Vilber Lourmat, Marne-la-Vallee, France). Densitometry was performed by ImageJ software.

4.4. Imaging of FAM AO Internalization. H2K mdx cells were cultured and differentiated as described previously. For internalization experiment, fluorescein (FAM)-labeled AOs were incubated at 400 and 800 nM concentrations in a final incubation volume of 500 μL/well in a 24-well plate. After 72 h of incubation, cell nuclei were stained with Hoechst for 10 min and washed five times with phosphate-buffered saline containing 10% fetal bovine serum before the images were captured using an Olympus TS-100 inverted fluorescence microscopy system.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/10.1021/acsomega.0c01463. RT-PCR and transcript densitometry analyses (PDF)

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Notes

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