Optimal sequence of antisense DNA to silence YB-1 in lung cancer by use of a novel polysaccharide drug delivery system

HIROTO IZUMI, SHOHEI NAGAO, SHINICHI MOCHIZUKI, NOBUAKI FUJIWARA, KAZUO SAKURAI and YASUO MORIMOTO

1University of Occupational and Environmental Health, Yahatanishi-ku, Kitakyushu, Fukuoka 807-8555; 2Department of Chemistry and Biochemistry, University of Kitakyushu, Wakamatsu-ku, Kitakyushu, Fukuoka 808-0135, Japan

Received August 15, 2015; Accepted December 16, 2015

DOI: 10.3892/ijo.2016.3451

Abstract. Silencing Y-box binding protein 1 (YB-1) can be an excellent target for cancer therapy and many lung cancer cells express the polysaccharide-recognition receptor Dectin-1. We designed a Dectin-1 targeting vehicle delivering YB-1-antisense DNA. First, we selected five optimal antisense DNA sequences to silence YB-1 from among 153 candidates. We chose the sequence closest to the start codon (AS014), and attached dA40 to the 3' end; dA40 promotes complex formation with a β-(1→3)-D-glucan called schizophyllan (SPG). The resultant complexes were applied to 12 human-oriented lung cancer cell lines, and cell viability was examined. The cell lines exhibited decreased viability and showed strong affinity to bind SPG, suggesting the AS014/SPG complex entered the cells via the Dectin-1 mediated pathway.

Introduction

Y-box binding protein 1 (YB-1) was originally found to initiate gene transcription (1). Its transcription is triggered by binding between YB-1 and inverted CCAAT box, which is normally located upstream of the TATA box in the promoter region. YB-1 has been shown to be involved not only in the transcription of various genes but also in cell viability and DNA repair (2). Furthermore, many studies demonstrate that YB-1 is overexpressed in human cancer cell lines; therefore, YB-1 may be a good target for cancer therapy (3). Shibahara et al (4) reported that most non-small lung cancers overexpress YB-1. Therefore, silencing YB-1 may also suppress the growth of non-small lung cancers.

Silencing the expression of a particular gene in cancer cells is critical in cancer therapy, especially when the gene in question is related to the growth and malignant alteration of cancer cells. Therapeutic oligonucleotides including antisense DNA and siRNA specifically silence protein expression by interacting with target mRNA; thus, they are presumably more efficient and less toxic than conventional low-molecular drugs (5,6). As the target mRNAs are located in the cytoplasm, it is essential for antisense DNA or siRNA to be transported to the cytoplasm in order to have an effect. Oligonucleotides themselves do not have the ability to be ingested by cells; furthermore, they are quite fragile in biological fluids because of enzymatic degradation and non-specific adsorption by serum proteins. Therefore, they require a drug delivery system (DDS) to protect them from such unfavorable interactions and transport them to the cytoplasm. The most commonly used synthetic DDS particles for oligonucleotide delivery are cationic lipids and polymers, which form electrostatic complexes with negatively charged oligonucleotides (7). Although some of these cationic compounds are used for cellular transfection in the laboratory, many problems must be overcome before they can be used in humans, including low uptake efficiency owing to a lack of targeting ability.

The natural polysaccharide schizophyllan (SPG) (Fig. 1) has a main chain comprising β-(1→3)-D-glucan and one β-(1→6)-D-glycosyl side chain that links to the main chain at every three glucose residues (8). We found that SPG forms a complex with single-stranded homo-polynucleotides and examined the fundamental properties of this complex (9-11). We recently started to apply this complex to therapeutic oligonucleotide delivery, including CpG DNA (12) and siRNA (13). For siRNA delivery, we demonstrated that the SPG/siRNA complex can be recognized by a polysaccharide receptor called Dectin-1 and subsequently enter the endocytic pathway (13,14). This indicates SPG-mediated siRNA delivery is one of the few known systems that possess cell-specific targeting. Heyl et al (15) demonstrated that Dectin-1 is widely expressed on human lung tissues. Therefore, there is a strong possibility that some of the lung cancers also express Dectin-1, meaning they could be treated by delivering antisense oligonucleotides (AS-ODNs) to silence YB-1.
Materials and methods

Preparation of antisense oligonucleotide sequence for YB-1. YB-1 mRNA consisting of 1,561 bases was obtained from the NCBI database (NCBI Reference Sequence: NM_004559.3) (16). To find an effective antisense sequence, we synthesized 153 different AS-ODNs in which the length of the base was 25 and each sequence was designed to match a part of the YB-1 mRNA sequence shifted from the 5' end 10 by 10 bases. The AS-ODNs were designated AS0019-AS153 (Fig. 2).

**SPG and complexation.** SPG (average molecular weight, 4.5x10⁵) was kindly provided by Mitsui Sugar Co., Ltd. (Tokyo, Japan); this is the same SPG used in other DDS studies (13). We prepared AS-ODNs/SPG complexes from SPG and phosphorothioate AS-ODNs with the (dA)₄₀ tail for YB-1. All phosphorothioate oligonucleotides including AS-ODNs and Alexa546-labeled dA₄₀ nucleotide were synthesized by Gene Design Co., Ltd (Osaka, Japan) and purified by high-performance liquid chromatography. SPG was dissolved in 0.25 N NaOH (aq) for 2-5 days to dissociate the triple helix into a single chain. The SPG solution, AS-ODNs in water, and phosphate-buffered solution (330 mM Na₂HPO₄, pH 4.7) were mixed at 37°C for 10 minutes to form the complex.
mixed, then the mixture (60 µM AS-ODNs, pH 7.4) was stored overnight at 4°C.

**Lung cancer cell lines.** The following 12 lung cancer cell lines were used: B203L, PC9, A110L, A549, HI299, QG56, SQ1, B1203L, PC10, 904L, PC1 and A529L. Their characteristics and other information are described elsewhere (17,18). All cell lines were cultured in RPMI-1640 medium, GlutaMAX™ (Life Technologies, Tokyo, Japan) and maintained in a 5% CO2 atmosphere at 37°C.

**Cell viability assessment by the water-soluble tetrazolium salt-8 (WST-8) assay.** The cell viability assay has been previously described (19). Briefly, PC9 cells (1x10^3) was seeded into 96-well plates. After 24 h, AS-ODNs were transiently transfected into the cells with 0.4 µl of RNAiMax (Life Technologies) per well. Final concentration of AS-ODNs is 50 nM in 120 µl of medium. For AS-ODNs/SPG complex, they were directly added into the medium. After 96 h, the surviving cells were stained with TetraColor One (Seikagaku Corp., Tokyo, Japan) for 2-3 h at 37°C according to the manufacturer’s instructions. The absorbance was then measured at 450 nm.

**Uptake of FITC-labeled SPG and Alexa546-labeled dA_40 nucleotide/SPG complex.** Cells were seeded at 2.5x10^4 cells/well in 24-well plates and incubated at 37°C under 5% CO2. The cells were cultured in RPMI-1640 containing 10% FBS and 100 U/ml penicillin and 0.1 mg/ml streptomycin. After 24 h, 0.1 µM FITC-labeled SPG (FITC-SPG) (13) or 0.5 µM SPG complexed with Alexa546-labeled dA_40 nucleotide (A546-dA_40/SPG) was added to cells in the presence of serum. After 8 h, cells treated with FITC-SPG were washed twice with PBS and was observed with EVOS® FL imaging system (Life Technologies). For competition assay, 10 µM (20 times excess) SPG complexed with unlabeled dA_40 nucleotide (dA_40/SPG complex) was used to treat the cells at the same time as the administration of 0.5 µM A546-dA_40/SPG. After 6 h, cells were washed twice with PBS and was observed with EVOS® FL imaging system.

**Western blot analysis.** PC9 cells were suspended in lysis buffer with 10 mM Tris-HCl (pH 7.9), 150 mM NaCl, 0.5% NP40, 1 mM PMSF and sonicated for 10 sec. Whole cell lysate (2.5, 5 and 10 µg) were separated on a 10% SDS-PAGE gel and transferred to a PVDF microporous membrane (Millipore, Bedford, MA, USA). The membrane was immunoblotted with anti-YB-1 antibody (20) at a 1:10,000 dilution, or with an anti-β-actin antibody (A5441; Sigma-Aldrich) at a 1:10,000 dilution, for 1 h, and then incubated with HRP-conjugated anti-rabbit IgG or anti-mouse IgG for 40 min. Detection was performed using enhanced chemiluminescence (GE Healthcare, Tokyo, Japan). Protein expression levels were quantitated using Multi Gauge version 3.0 (Fujifilm, Tokyo, Japan).

**Results and Discussion**

**Selecting the antisense sequence.** The cell viability of cancer cells treated with all AS-ODNs transfected into PC9 cells is shown in Fig. 3; this is an *in vitro* screen for cell viability. We used the RNAiMax reagent, a Lipofectamine transfection reagent, which is a cationic liposome formulation that forms positively charged polyion complexes with negatively charged DNA or siRNA. Such complexes allow the bound DNA or siRNA to eventually cross into the cytoplasm. As YB-1 is overexpressed in PC9 cells (4) the observed decreased cell viability is most likely due to YB-1 silencing by AS-ODNs; moreover, the differences in cell viability reflect the efficacy of individual AS-ODNs. The five most effective sequences are indicated by arrows. Four AS-ODNs-AS010, AS014, AS025, and AS030, which decreased the cell viability below one third are close to the start codon, while AS101 is not (Fig. 4).
The recruitment of RNase H1 to the antisense DNA/mRNA duplex is widely believed to be the key step for protein silencing. However, there are few such sites on mRNAs, because the binding site of mRNA must be a single chain and exposed to the outside to make it easy for antisense DNA to bind the site. The bulge/loop structure of YB-1 mRNA was estimated using UNAfold (21); the result is shown in Fig. 4, and the binding sites of the four AS-ODNs are indicated by solid lines.

![Figure 4. The mRNA bulge/loop structure of YB-1 generated with UNAfold, and the binding sites of the selected five AS-ODNs.](image)

The YB-1 protein expression was silenced in PC9 owing to antisense DNA, determined by western blot analysis. (A) PC9 cells were transfected with control oligonucleotide (Ctrl) and indicated AS-ODNs. After 72 h, whole cell lysate (10, 5, and 2.5 µg) were subjected to western blot analysis with anti-YB-1 or anti-β-actin antibody. (B) Each expression level of YB-1 was normalized to that of β-actin using signal intensity of 5 or 2.5 µg cell lysate and compared the YB-1 expression level with ctrl mock transfection. The ratio of relative YB-1 expression is shown in the bottom. Each control expression was set to 1.

![Figure 5. YB-1 protein expression was silenced in PC9 owing to antisense DNA, determined by western blot analysis.](image)

The recruitment of RNase H1 to the antisense DNA/mRNA duplex is widely believed to be the key step for protein silencing. However, there are few such sites on mRNAs, because the binding site of mRNA must be a single chain and exposed to the outside to make it easy for antisense DNA to bind the site. The bulge/loop structure of YB-1 mRNA was estimated using UNAfold (21); the result is shown in Fig. 4, and the binding sites of the four AS-ODNs are indicated by solid lines. All
AS-ODNs binding sites contain loop structures. Therefore, these loops are hypothesized to be responsible for the YB-1 silencing; thus, after hybridization, the double DNA/mRNA strand is presumably recruited and cleaved by RNase H1.

The results of western blot analysis comparing the expression of five AS-ODNs silencing YB-1 are shown in Fig. 5A. Four AS-ODNs (AS10, AS14, AS25 and AS30) attenuated YB-1 expression to <50% of the control (Fig. 5B). These results confirm that the decreased cell viability is due to YB-1 silencing after transfection. Western blot analysis indicated that AS014 and AS025 reduced the YB-1 expression very efficiently. As AS014 is closest to the start codon region and exhibited considerably good efficacy in both assays, we used AS014 hereafter.

Targeted delivery of AS-ODN with SPG complexation. SPG forms a stoichiometric complex with particular homonucleotides such as poly(C) or poly(dA) via combination of hydrogen bonding and hydrophobic interactions (9-11). On the basis of our previous studies, we attached dA40 to the AS-ODN to enable complex formation with SPG and thus efficient gene silencing (13,14). When the attachment position is the 3' end of the AS-ODNs, the phosphorothioate linkage forms more stable complexes than the phosphodiester ones (13). Therefore, we attached phosphorothioate dA40 to the 3' end of AS014 to form a complex with SPG. The exact stoichiometric composition is (mG):(dA)=2:1, where mG is the main chain glucose (9,10). However, in practice, we normally prepare the complexes at an SPG rich composition. In the present assay, we prepared the AS014-dA40/SPG complex at (mG):(dA)=4:1. No free AS014-dA40 was observed in gel electrophoresis (Fig. 6).

We previously demonstrated that the complex is taken up by Dectin-1 expressing immunocytes. We recently cloned and purified the extracellular domain of murine Dectin-1 and performed binding affinity analysis between this protein and the SPG/DNA complex by using quartz crystal microbalance (22). Phosphorothioate dA 40 markedly enhanced the Dectin-1 binding affinity compared to that with phosphodiester and also behaved differently in the Dectin-1 mutants in which the Trp221 and His223 residues in the 3'-terminal exon were replaced with alanine. There appeared to be multiple binding sites: the same site as SPG and an additional site(s) in which phosphate anion specific electrostatic interactions were mainly involved. This enhanced affinity of the phosphorothioate DNA/SPG complex is another reason prompting its use in this experiment.
The cell viability rates of various lung cancer cells after the AS014/SPG complex was applied at 0.4 or 1.0 µM are shown in Fig. 7. There are over 200 known lung cancer cell lines, from small to non-small cell types, exceeding the number of other common epithelial cancers. Among them, we selected 12 commonly available cell lines. AS014/SPG complex application resulted in variably decreased cell viability rates. As all of these cells overexpress YB-1, which was also confirmed in the present study (data not shown), the decreased cell viability is mainly attributable to the number of AS014 molecules ingested and bound to YB-1. However, it is unknown whether these cells express Dectin-1; this issue is currently being investigated. The greatest decrease in cell viability was observed in B203L, PC9, B1203L and PC10 cells (Fig. 7A). AS014/SPG complex decreased YB-1 expression ~40% in PC9 cells, but not A549 cells (Fig. 7B). This results are consistent with cell viability assessment (Fig. 7A). We selected B203L and PC9 as well as A549 and PC1 cells, which showed no decrease in cell viability, and performed fluorescent microscopy to observe FITC-labeled SPG. SPG was actively ingested by B203L and PC9 but not A549 or PC1 cells (Fig. 8A), which is concordant with their cell viability. We also found A546-dA40/SPG was ingested by PC9 cells (Fig. 8B). This ingestion was partially abolished by addition of the dA40/SPG complex (Fig. 8C). As mentioned above, the Dectin-1 expression assay for these cells is ongoing. Nevertheless, it is safe to conclude that AS014 was ingested by B203L and PC9 cells, which was facilitated by the recognition of Dectin-1 for the AS014/SPG complex. Furthermore, the results indicate AS014 silenced the YB-1 expression, resulting in decreased cell viability.

In conclusion, five optimal antisense DNA sequences for silencing YB-1 expression in lung cancers were selected from among 153 candidates. We chose the one closest to the start codon, AS014, and attached dA40 to the 3' end to form a complex with SPG. The resultant complexes were applied to 12 human-oriented lung cancer cell lines to examine cell viability. Several cell lines exhibited markedly decreased cell viability presumably due to YB-1 silencing, which was corroborated by western blot analysis. The cells exhibited such decrease also ingested SPG, suggesting the AS014/SPG complex entered the cells via the Dectin-1 mediated pathway.
References


