In vitro optimization of 2'-OMe-4'-thioribonucleoside–modified anti-microRNA oligonucleotides and its targeting delivery to mouse liver using a liposomal nanoparticle

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Received July 14, 2013; Revised August 20, 2013; Accepted August 21, 2013

ABSTRACT

MicroRNAs (miRNAs) are small noncoding RNAs that regulate gene expression post-transcriptionally. Previous studies, which characterized miRNA function, revealed their involvement in fundamental biological processes. Importantly, miRNA expression is deregulated in many human diseases. Specific inhibition of miRNAs using chemically modified anti-miRNA oligonucleotides (AMOs) can be a potential therapeutic strategy for diseases in which a specific miRNA is overexpressed. 2'-O-Methyl (2'-OMe)-4'-thioRNA is a hybrid type of chemically modified oligonucleotide, exhibiting high binding affinity to complementary RNAs and high resistance to nuclease degradation. Here, we evaluate 2'-OMe-4'-thioribonucleosides for chemical modification on AMOs. Optimization of the modification pattern using a variety of chemically modified AMOs that are perfectly complementary to mature miR-21 revealed that the uniformly 2'-OMe-4'-thioribonucleoside–modified AMO was most potent. Further investigation showed that phosphorothioate modification contributed to long-term miR-122 inhibition by the 2'-OMe-4'-thioribonucleoside–modified AMO. Moreover, systemically administrated AMOs to mouse using a liposomal delivery system, YSK05-MEND, showed delivery to the liver and efficient inhibition of miR-122 activity at a low dose in vivo.

INTRODUCTION

MicroRNAs (miRNAs) are a class of endogenously expressed small noncoding RNAs (18–25 nt), which regulate gene expression post-transcriptionally. In animals, single-stranded mature miRNAs hybridize to the 3' untranslated region (3'UTR) of the target mRNA through complete base paring with positions 2–8 of the miRNA, known as the seed region. Binding on the seed region nucleates miRNA–mRNA association, and causes translational inhibition or mRNA degradation (1,2). Because complementarity of the seed region consists of only 7 nt, a single miRNA may regulate multiple genes, and a single mRNA can be modulated by several different miRNAs (3–5). To date, >1000 miRNAs have been identified in humans and regulate up to 60% of protein coding genes (6,7). MiRNAs are implicated in important functions in the biological process, including cell differentiation, proliferation, development, metabolism and apoptosis. Furthermore, up- or downregulation of miRNA expression is correlated with a variety of human diseases such as cancer, viral infection and cardiovascular disorders (8). Thus, regulation of specific miRNA function is a promising therapeutic strategy for treatment of such diseases.

Among the approaches to modulate the function of miRNAs, anti-miRNA oligonucleotide (AMO)-based inhibition has been most widely used not only to exploit the biological function of miRNAs but also as candidates for therapeutic agents (9). To develop oligonucleotide (ON)-based therapeutic strategies, there are several issues to overcome, namely poor stability of ONs in biological fluids, weak binding affinity to target RNA, poor cellular uptake and unfavorable immunostimulatory activity. Thus far, a wide variety of chemically modified ONs have been developed to date, including 2'-O-methyl (2'-OMe) (10–13), 2'-O-methoxymethyl (14) and locked nucleic acid (LNA) to overcome these disadvantages (15–17). These chemical modifications have successfully been applied to an antisense technology as well as...
ON-based therapeutic technologies (e.g. siRNAs, aptamers, ribozymes). Two successful reports of 2'-OMe–modified AMOs were reported in 2004 (18,19), lending credence to the idea of using chemically modified AMOs. Furthermore, considerable efforts in optimization have been dedicated to develop AMOs as a new therapeutic agent, and several chemically modified AMOs are currently undergoing clinical trials (20). Therefore, further optimization of chemical modifications for AMO-based miRNA suppression will continue to improve this therapeutic approach.

We developed a novel chemically modified ON, 2'-OMe-4'-thioRNA (21) (Figure 1), which can be considered a hybrid chemical modification based on 2'-OMe RNA (13) and 4'-thioRNA (22–28). In our previous study, we reported the development of 2'-OMe-4'-thioribonucleoside–modified siRNA (29). Optimization of both the number and position of modification by 2'-OMe-4'-thioribonucleoside afforded modified siRNAs with more potent and persistent RNAi activity. In addition, investigation of the duration of RNAi activity resulted in long-lasting gene silencing in vitro owing to the improved intracellular stability of 2'-OMe-4'-thioribonucleoside–modified siRNA. Like many other chemically modified ODNs that can successfully be applied to AMO as well as siRNA, we expected that 2'-OMe-4'-thioribonucleoside modification acts as a promising AMO. Therefore, we set out to evaluate the utility of 2'-OMe-4'-thioribonucleoside for chemical modification on AMOs. In this study, we investigated the modification pattern of AMOs by 2'-OMe-4'-thioribonucleoside in terms of potency and duration of activity in two kinds of target miRNA (miR-21 and miR-122) in vitro. Moreover, systemically administrated AMOs to mouse liver using a liposomal delivery system, a multifunctional envelope-type nano device (MEND) with a pH-sensitive cationic lipid, YSK05 (YSK05-MEND) (30), showed efficient inhibition of miR-122 activity at a low dose in vivo, implicating potential use of 2'-OMe-4'-thioribonucleoside–modified AMO in nucleic acid therapy.

**MATERIALS AND METHODS**

**Oligonucleotides**

The chemically modified AMOs used in this study were synthesized on an Applied Biosystem 3400 DNA synthesizer according to our previous report (21). Thus, support bound chemically modified AMOs were synthesized using the corresponding phosphoramidite units at a 1.0 μmol scale following the standard procedure described for oligoribonucleotides. Each of the phosphoramidite units was used at a concentration of 0.1 M in dry acetonitrile, and the coupling time was extended to 10 min for each step. AMOs with phosphorothioate (PS) backbone were achieved by oxidation with 3H-1,2-benzodithiol-3-one-1,1-dioxide (Beaucage reagent) during ON synthesis. After completion of the synthesis, the CPG support was treated with concentrated NH₄OH or NH₄OH/EtOH (3:1) at 55°C for 16 h. In the case of CPG supports containing either 2'-F or 2'-F-4'-thioribonucleoside modification, these were treated with methanolic ammonia (saturated at 0°C) at room temperature for 24 h. Then, the support was filtered off. The filtrate was concentrated and the ON protected by a DMTTr group at the 5'-end was chromatographed on a C-18 silica gel column with a linear gradient of acetonitrile (from 5 to 40%) in 0.1 N TEAA buffer (pH 7.0). The fractions containing the full-length ON were combined and concentrated. The residue was treated with aqueous acetic acid (70%) for 20 min at room temperature. The solution was concentrated and the residue was purified on reversed-phase high performance liquid chromatography, using a J’sphere ODS-M80 column (4.6 × 150 mm, YMC) with a linear gradient of acetonitrile (from 10 to 40%) in 0.1 N TEAA buffer (pH 7.0). The structures of each RNA were confirmed by measurement of MALDI-TOF/MASS spectrometry on Ultraflex TOF/TOF (Bruker Daltonics). The analytical data of synthetic AMOs are summarized in Supplementary Table S1.

**Tₘ measurement**

Thermally induced transitions were monitored at 260 nm on a Beckman DU 650 spectrophotometer. Samples were prepared as follows: AMO and target miRNA (3 μM each) were mixed in a phosphate buffer (10 mM, pH 7.0) containing 0.1 mM EDTA and 1 mM NaCl, heated at 90°C for 5 min, cooled gradually to room temperature and used in thermal denaturation studies. The sample temperature was increased 0.5°C/min.

**Cell culture**

HeLa cells and Huh-7 cells were cultured at 37°C under 5% CO₂ in Dulbecco’s modified Eagle’s medium (DMEM, Gibco) supplemented with 10% fetal bovine serum (FBS, Thermo Fisher Scientific), 100 units ml⁻¹ penicillin, 100 μg ml⁻¹ streptomycin, respectively. Cells were regularly passaged to maintain exponential growth.

**Construction of miRNA reporter plasmids and luciferase reporter assay**

The miRNA reporter plasmids were generated by cloning the synthetic 5’-phosphorylated ONs corresponding to tandem perfect-match target sited for human miR-21 or miR-122 into the 5’ UTR of firefly luciferase gene (luc2) in the pmirGLO Vector (Promega). HeLa cells or Huh-7 cells were plated into 96-well plates, at 10 000 cells/well in DMEM supplemented with 10% FBS, 100 units ml⁻¹ penicillin, 100 μg ml⁻¹ streptomycin. After 24 h of plating, reporter plasmid and AMOs were co-transfected into the cells in triplicate using Lipofectamine™ 2000 transfection reagent.

**Figure 1. Structure of 2'-modified-4'-thioribonucleoside.**
(Invitrogen) according to the manufacturer’s instructions in 100 µl/well Opti-MEM (Invitrogen). AMO concentrations varied as indicated, whereas plasmid concentrations remained constant at 0.1 µg/well. After 6 h of transfection, cells were re-fed with complete media. Cells were lysed at indicated hours after transfection, and luciferase activities were measured using the Dual-Luciferase Reporter Assay System (Promega). Relative firefly luciferase signals were normalized against Renilla luciferase signals. Results are expressed as relative Fluc/Rluc ratios to that of mirGLO-treated cells. Each experiment was performed at least three times.

**Preparation of AMO encapsulated liposome**

As a liposomal delivery system, YSK05-MENDs encapsulating AMOs were prepared by a t-BuOH dilution procedure (30,31). Thus, each AMO was mixed with 90% (v/v) t-BuOH containing YSK05 (synthesized as previously described) (30), cholesterol (Avanti Polar Lipid), 1,2-dimyristoyl-sn-glycerol and methoxyethyleneglycol 2000 ether (Avanti Polar Lipid) at a molar ratio of 70:30:3 in 20 mM citrate buffer (pH 4.0) at AMO/lipids of 0.1 (wt/wt) under strong agitation to a t-BuOH concentration of 60% (v/v). Then, lipids/AMO mixture was added into 20 mM citrate buffer (pH 4.0) under strong agitation to a t-BuOH concentration of <12% (v/v). Ultrafiltration was performed to remove t-BuOH, replace external buffer with phosphate buffered saline (PBS, pH 7.0) and concentrate a resulting YSK05-MEND encapsulating AMO (AMO-YSK05-MEND). The average diameter and zeta-potential of the AMO-YSK05-MENDs were determined using a Zetasizer Nano ZS ZEN3600 (MALVERN Instrument, Worcestershire).

**In vivo experiments**

Female BALB/c mice (8 weeks old) were purchased from Japan SLC. All in vivo experiments were approved by the Institutional Animal Care and Use Committee. One day before the administration, serum was collected for measuring cholesterol concentration. Each AMO-YSK05-MEND was diluted to the appropriate concentrations in PBS (pH 7.4), followed by intravenous administration via the tail vein at a dose of 1 mg AMO/kg in 10–15 ml/kg given once a day, every other day, for three times. At 48 h after the last injection, liver and blood were collected. Blood sample was centrifuged at 8g at 4°C for 5 min to obtain plasma. To obtain serum, blood sample was stored overnight at 4°C, followed by centrifugation (10,000 rpm, 4°C, 10 min). Cholesterol concentration in plasma and alanine aminotransferase level in serum were determined using a cholesterol E-test WAKO and Transaminase CII-test WAKO (Wako) according to the manufacturer’s protocols. Total RNAs in liver were isolated using TRizol (Invitrogen) according to the manufacturer’s protocols. Then, isolated RNAs were reverse transcribed using a High Capacity RNA-to-cDNA kit (Applied Biosystems) according to manufacturer’s protocol. A quantitative polymerase chain reaction (PCR) analysis was performed with 20 ng of cDNA using Fast SYBR Green Master Mix (Applied Biosystems) on the Lightcycler480 System II (Roche Applied Science). All reactions were performed at a volume of 15 µl. The primers for qRT-PCR are as follows: mouse AldoA (forward) 5′-GATGGGTCCAGC TTCAAC-3′ and (reverse) 5′-GTGCTTCTTCTCTTCCA A CTTG-3′; mouse Bckdk (forward) 5′-AGGACCTATG CATGGCTTTG-3′ and (reverse) 5′-CCGTAGGTAGAC ATCCGTTG-3′; mouse Ndrg3 (forward) 5′-ATGGGCTA CATACCATCTG-3′ and (reverse) 5′-TCTGACTGATT GCTGTGAC-3′; mouse Hprt1 (forward) 5′-CGTGATT AGGGATGAGAAC-3′ and (reverse) 5′-GCAAGTCTTTC AGATCTGAC-3′. Each data was normalized by Hprt1 expression. All experiments were performed in triplicate and data show mean values from at least three assays.

**Statistical analysis**

Comparisons between multiple treatments were made using one-way analysis of variance (ANOVA), followed by the SNK test. Pair-wise comparisons between treatments were made using a student’s t-test. P < 0.05 was considered significantly different.

**RESULTS AND DISCUSSION**

**Evaluation of inhibitory activity of AMOs against miR-21**

We first synthesized modified AMOs against miR-21, a miRNA that is overexpressed in many tumors and thus is considered to be a potential therapeutic target in oncology (32). The sequences and modification patterns of AMOs were shown in Table 1. Because 2′-F-4′-thioRNA showed highest hybridization ability with its complementary RNA among the chemically modified ONs tested (21,33), we synthesized chimeric 2′-OMe-4′-thio-2′-F-4′-thio–modified AMOs (AM21SMF1 and AM21SMF2) along with 2′-OMe-4′-thio–modified AMO (AM21SM), which are complementary and the same length (22-mer) as mature miR-21. For comparison, 4′-oxy congeners (AM21M, AM21MF1 and AM21MF2) were also prepared. Because Hutvagner et al. reported that AMOs possessing a complementary sequence core with 5′nt flanking sites on both 5′- and 3′-ends showed more potent anti-miRNA activity (19), we also designed a 32-mer uniformly 2′-OMe-4′-thioribonucleoside–modified AMO (AM21SM-L), which exhibits perfect complementarity to miR-21 and 5 nt flanking regions on both ends, as well as that of 2′-OMe (AM21M-L).

To evaluate anti-miRNA activity of the synthetic AMOs, we constructed a miR-21 luciferase reporter plasmid (34), which has two perfectly matched miR-21 binding sites arranged in tandem on 3′ UTR of firefly luciferase gene (pmirGLO21). The firefly luciferase activity was completely suppressed (3.7%) when the pmirGLO21 was transfected into HeLa cells, a cell line known to have high endogenous levels of miR-21 (Figure 2). The activity of pmirGLO, which has no miR-21 binding sites, was high (normalized as 100%) on transfection into HeLa cells. Our hypothesis is that introduction of miR-21 AMO can prevent miR-21 binding to the pmirGLO21 target sites, resulting in increased
dependent manner. In general, our AMOs showed miR-21 inhibitory activity in a dose-

Figure 2. Anti-miR-21 activity of modified AMOs. HeLa cells were co-transfected with the miR-21 reporter plasmid (mirGLO21) and the indicated modified anti-miR-21 AMOs at the indicated concentrations. The dual luciferase assay was performed at 24 h after transfection. Data are shown as mean with standard deviation (SD).

Table 1. Sequence and modification patterns of anti-miR-21 AMOs

| AMO       | Sequencea | Tm (°C)b |
|-----------|-----------|----------|
| AM21SM    | 5'-UCAAACAUCAGCUGUAAGCUA-3' | 64.1 |
| AM21SMF1  | 5'-AcAcAcAcAcAGCuAcAgCuA-3' | 69.2 |
| AM21SMF2  | 5'-UCAAACAUCAGCUGUAAGCUA-3' | 71.1 |
| AM21M     | 5'-UCAAACAUCAGCUGUAAGCUA-3' | 59.9 |
| AM21MF1   | 5'-AcAcAcAcAcAGCuAcAgCuA-3' | 64.2 |
| AM21MF2   | 5'-UCAAACAUCAGCUGUAAGCUA-3' | 67.9 |
| AM21SM-L  | 5'-UCUUAACAAACAUCAGCUGUAAGCUAACCUU-3' | 62.1 |
| AM21M-L   | 5'-UCUUAACAAACAUCAGCUGUAAGCUAACCUU-3' | 58.0 |

aUppercase letters represent 2'-OMe; lowercase letters represent 2'-F; bold uppercase letters are 2'-OMe-4'-thioribonucleotides; bold lowercase letters are 2'-F-4'-thioribonucleotides. bTm values were measured versus with target miR-21 (22-mer) in a phosphate buffer (10 mM, pH 7.0) containing 0.1 mM EDTA and 1 mM NaCl, 3 μM strand concentration. Values were given as an average of three independent experiments.

luciferase expression by either causing translational inhibition or cleavage of the mRNA. We then tested anti-

It is known that there is a good correlation between Tm value of AMOs and in vitro potency (35). Therefore, we compared the thermal stability of the AMOs with target miR-21 based on their Tm values. As can be seen in Table 1, the Tm values of chimera AMOs, which contain either 2'-F or 2'-F-4'-thioribonucleoside modification, were higher than those of the uniformly modified AMOs. Although AM21SMF2 showed the highest Tm value, its anti-miR-21 activity was lowest among 4'-thio-modified AMOs. A similar trend was observed in a series of AMOs consisting of 4'-oxo congeners. Thus, no correlation was observed between activity and binding affinity in our case. Meanwhile, as we reported previously (21), 2'-OMe-4'-thioribonuclease showed higher nuclease stability compared with 2'-OMe RNA, 2'-F-4'-thioribonuclease and 2'-F RNA. Therefore, chimera AMOs might be more susceptible to degradation than the uniformly modified AMOs, serving as a potential explanation why AM21SM showed the highest activity amongst all AMOs tested. From these results, it can be concluded that the 2'-OMe-4'-thioribonucleoside modification is useful for inhibition of miRNA function. Also, uniform modification is simple and seems to be the most promising choice for subsequent studies.

Improvement in duration of anti-miRNA activity of AMOs by backbone substitution

To examine if 2'-OMe-4'-thioribonucleoside–modified AMO is capable of inhibiting other miRNAs, we chose miR-122 as another miRNA target. MiR-122 is a liverspecific miRNA, known to be involved in cholesterol metabolism, fatty acid metabolism (36) and hepatitis C virus replication (37). Many successful studies of miR-122 inhibition by AMOs both in vitro and in vivo have been reported (20,36-42), and some of them are currently under investigation in clinical trials (20). We hypothesized that 2'-OMe-4'-thioribonucleoside–modified AMO could have high potency to inhibit miR-122 as well as miR-21.

We synthesized a uniformly 2'-OMe-4'-thioribonucleoside–modified AMO, which is complementary to and the same length (23-mer) as mature miR-122,
with unmodified phosphodiester (PO) backbone (AM122SM, Table 2). Because ONs can be degraded by nuclease by cleaving a phosphodiester linkage in biological fluid, substitution of a PO linkage for a PS linkage is ideal to prevent such nuclease cleavage reactions. In fact, PS modification has successfully been applied to many previously reported AMOs. Because our end goal of this study is in vivo application, we synthesized 2′-OMe-thioribonucleoside–modified AMOs with PS backbones (AM122SM-PS). For comparison, we also synthesized uniformly 2′-OMe–modified AMOs with either PO or PS backbones (AM122M and AM122M-PS, respectively). Recently, Obad et al. developed seed targeting 8-mer tiny LNAs, which can simultaneously inhibit miRNA families that share the same seed region (42). According to this report, we also prepared a seed targeting 8-mer (AM122SM-PS 8 nt) along with two shorter sequences (AM122SM-PS 15 nt and AM122SM-PS 20 nt) to examine their potency as well as to find an optimal length of AMOs.

Each AMO and a miR-122 reporter plasmid (pmirGLO122), constructed in a same manner to pmirGLO21, were co-transfected into Huh-7 cells at various AMO concentrations (0.5–5 nM). At 24 h after transfection, we harvested the cells and carried out dual luciferase reporter assay. The resulting AMO activities were shown in Figure 3. As was the case of miR-21 inhibition, all AMOs showed miR-122 inhibitory activity in a dose-dependent manner, and AM122SM gave higher activity than AM122M (63.9 versus 36.6% at 5 nM). No obvious difference was observed in the activity between AMOs with PO and PS backbones. Concerning of the length of AMO, anti-miRNA activity decreased as AMO length became shorter, and tiny AMO, namely AM122SM PS 8 nt, did not show any activity. Thus, the 2′-OMe-4′-thioribonucleoside–modified AMO possessing complementary matched sequence and length seems to be suitable for miRNA inhibition.

In our previous study, we showed prolonged activity of 2′-OMe-4′-thioribonucleoside–modified siRNA (29). Thus, we expected that the 2′-OMe-4′-thioribonucleoside modification can prolong the activity of AMOs. We next performed a time-course experiment. As described above, no difference between PO and PS AMOs was observed in the activity after 24 h. However, considerable differences were observed after 48 h, where the activities of PO AMOs (AM122SM and AM122M) declined (Figure 4), while activities of PS AMOs (AM122SM-PS and AM122M-PS) increased. AM122SM-PS showed dramatic increase of the activity throughout the assay, as much as 90% at 72 h. From these results, it can be concluded that 2′-OMe-4′-thioribonucleoside–modified AMO are superior compared with 2′-OMe–modified AMO, and that PS modification conferred long-term activity on AM122SM.

In the case of the miR-21 study described above, we suggested that nuclease resistance is more closely related to AMO potency than the $T_m$ values. We wished to explore further which of these two factors correlated more strongly with the potency. To explore a correlation between the potency and physical properties of 2′-OMe-4′-thioribonucleotide–modified miR-122 AMOs in more detail, we first measured the $T_m$ values of duplexes with a target miR-122. As can be seen in Table 2, AM122M

### Table 2. Sequence and modification pattern of anti-miR-122 AMOs

| AMO         | Sequencea | $T_m$ (°C)b |
|-------------|------------|-------------|
| AM122SM     | 5′-ACAAACACAUUGUCACACUCCA-3′ | 65.8        |
| AM122SM-PS  | 5′-ACAAACACAUUGUCACACUCCA-3′ | 64.8        |
| AM122SM-PS 20 nt | 5′-ACAAACACAUUGUCACACUCC-3′ | 60.7        |
| AM122SM-PS 15 nt | 5′-CAACACUCAUUGUCACACUUCC-3′ | 54.6        |
| AM122M      | 5′-ACAAACACAUUGUCACACUCCA-3′ | 36.3        |
| AM122M-PS   | 5′-ACAAACACAUUGUCACACUCCA-3′ | 38.0        |

aUpper case letters represent 2′-OMe; bold upper case letters are 2′-OMe-4′-thioribonucleotides; underlined are PS backbone modification. b$T_m$s were measured versus miR-122 in a phosphate buffer (10 mM, pH 7.0) containing 0.1 mM EDTA and 1 mM NaCl, 3 μM strand concentration. Values were given as an average of three independent experiments.

Figure 3. Anti-miR-122 activity of the modified AMOs. Huh-7 cells were co-transfected with the miR-122 reporter plasmid (miGLO122) and the anti-miR-122 AMOs at the indicated concentrations. The dual luciferase assay was performed at 24 h after transfection. Data are shown as mean with SD.
had a $T_m$ value of 62.9°C and that of AM122M-PS was lower ($T_m = 58.0$°C and $\Delta T_m = 4.9$°C). On the other hand, AM122SM showed a higher $T_m$ value (65.8°C) compared with that of AM122M, and AM122SM-PS had the $T_m$ value of 64.8°C with only a slight decrease from that of AM122SM ($\Delta T_m = 1.0$°C). It is well known that the PS modification lowers the $T_m$ of ONs (~0.5–0.7°C per modification) (43,44). This decrease of the binding affinity would cause loss of inhibitory activity of AMOs. However, replacement of the PO backbone of AM122SM with a PS had no effect on the binding affinity of the AMO, a possible explanation for the retained AMO activity.

We then investigated the nuclease stability of the modified AMOs. A series of AMOs were labeled at the 5'-end with $^{32}$P and incubated in 50% human plasma. The reactions were analyzed by denaturing PAGE (Supplementary Figure S1). As expected, PS AMOs exhibited higher stability than that of PO AMOs, and as we reported previously, 2'-OMe-4'-thioribonucleoside–modified AMOs were slightly more stable than that of the corresponding 2'-OMe–modified AMOs. It is worth noting that AM122SM-PS was totally intact under these conditions, while AM122M-PS showed some mild degradation. The rank order of stability in plasma was AM122SM-PS > AM122M-PS > AM122SM > AM122M, indicating that AM122SM-PS is extremely stable against nuclease degradation in human plasma. These results suggested that the $T_m$ value of the AMO had relatively little impact on AMO potency, whereas nuclease resistance seemed to have much greater effect.

For therapeutic use, nuclease resistance in biological fluid, tissues and cells is a prerequisite feature of chemically modified ONs. Our previous comprehensive study of nuclease stability revealed that 2'-OMe-4'-thioribonucleoside was significantly stable against both exo- and endonucleases in spite of consisting of PO backbones (21). Here, we showed PS modification on the 2'-OMe-4'-thioribonucleoside–modified AMO can further improve its nuclease resistance. As a more specific scenario, the endonuclease activity of Ago2 cleaves the unselected strand of RNA duplexes loading into the RISC. It is thought that AMOs act primarily by hybridizing with mature miRNAs that have been loaded into the RISC. Wang et al. explained the molecular mechanisms of target RNA cleavage by a crystal structure of Ago protein (45). They found that both 2'-OMe and PS substitution at the cleavage site (positions 10'-11') disrupted Ago slicer activity, owing to configurations of the sulfur atoms in Ago protein. Thus, we suggest that the potency and long-term activity of the AM122SM-PS would be due in part to its resistance to cleavage by Ago. Also, PS modification could increase intracellular stability of AMOs, consequently anti-miRNA activity AMOs increased over time (Figure 4). Taken together, combinatorial use of 2'-OMe-4’-thioribonucleosides and a PS backbone is an attractive choice for therapeutic AMOs.

**Targeted delivery and anti-miR-122 activity of 2'-OMe-4'-thioribonucleoside–modified AMO in mouse liver**

*In vitro* studies showed that AM122SM-PS possessed the most favorable properties. Hence, we next carried out *in vivo* studies to assess whether AM122SM-PS could also inhibit miR-122 in mice.

Two aspects of successful nucleic acid therapeutics are delivery, followed by ON stability. We have developed a new liposomal nucleic acid delivery system, YSK05-MEND, which contains a pH-sensitive cationic lipid for efficient release of siRNAs from the endosome into the cytoplasm (29). Thus, we prepared YSK05-MEND for *in vivo* delivery of AMOs to the liver. YSK05-MENDs encapsulating AM122M-PS and AM122SM-PS represented comparable size, charge, polydispersity and encapsulation efficiency (Supplementary Table S2). We assessed *in vivo* efficacy of the modified miR-122 AMOs by treating mice three times with intravenous injection of 1 mg/kg YSK05-MEND formulated AM122SM-PS or AMO122M-PS every 2 days.

As described above, a single miRNA may control the levels of multiple mRNAs (3–6). MiR-122 is no exception, as many mRNAs whose expressions are directly controlled by miR-122 have been identified. Elmén et al. demonstrated inhibition of miR-122 using LNA-modified AMOs, LNA-antimiR (41). In their experiments, derepressions of four mRNAs, AldoA, Bckdk, NDRg3 and Cld320, all of which are direct targets of miR-122 in the mouse liver, were observed after treatment of mice with LNA-antimiR. Therefore, we conducted expression analysis of three miR-122 target mRNAs in the mouse liver, AldoA, Bckdk and Ndrg3 (Figure 5A), by real-time PCR 48 h after the last injection. The levels of all three mRNAs were higher in the AMO-treated mice compared with those treated with saline. It is worth noting that AM122SM-PS induced higher mRNA expression levels than AM122M-PS in all three mRNAs examined. We also observed statistically significant differences between AM122SM-PS and AM122M-PS in the expressions of both AldoA and Bckdk (P < 0.05).

We next examined the change in plasma cholesterol level associated with the increase in expression of these
mRNAs. There was no obvious difference between the AMO- and saline-treated mice at 1 day after the last dose, while drastic changes were observed at day 6, where the serum cholesterol levels were reduced by 57.7% for AM122SM-PS and 52.5% for AM122M-PS (Figure 5B) without any hepatotoxicity (Supplementary Figure S2). These results suggest that AM122SM-PS was properly delivered to the mouse liver by YSK05-MEND and elicited an anti-miR-122 effect.

In the previous reports (20,36–38,40–42), AMOs have been administered either as naked/saline formulated or as small molecule conjugates (e.g. cholesterol, a cell-penetrating peptide) to assist in vivo delivery. These approaches often required relatively high doses (ranging from 10 to 80 mg/kg for mice). Instead, our YSK05-MEND dosing strategy achieves efficient miR-122 AMO delivery to the liver. Although the mouse strains used for our experiments versus other researchers’ previous studies were different (e.g. we used inbred strain mice, BALB/c, which are considered to be genetically identical, whereas others used outbred strains, such as NMRI), we successfully increased miR-122 target mRNA expression, followed by a decrease in serum cholesterol level with three intravenous doses of 1 mg/kg AM122SM-PS in mice.

CONCLUSION
We demonstrated the inhibition of miRNAs with 2'-OMe-4'-thioribonucleoside–modified AMOs in vitro as well as in vivo. We first evaluated a variety of chemically modified AMOs that are complementary to mature miR-21, and found the potency of the uniformly 2'-OMe-4'-thioribonucleoside–modified AMO to be the
best in our series. Further investigation revealed that PS modification contributes to long-term miR-122 inhibition by the 2'-OMe-4'-thioribonucleoside–modified AMO.

For in vivo studies, we took advantage of an efficient delivery technology, YSK05-MEND, and successfully increased three miR-122 target mRNA levels in the AMO-treated mice liver, followed by a decline of serum AST. Although many successful in vivo AMO studies have been reported so far, to our knowledge, none of them have used liposome-like delivery systems. In our previous study, we confirmed that YSK05-MEND efficiently releases siRNAs into the cytoplasm in a pH-dependent manner (30). It is with this property that we believe we were able to achieve efficient AMO delivery in mice.

Together, here we showed not only the potency of 2'-OMe-4'-thioribonucleoside–modified AMOs but also the utility of YSK05-MEND for AMO delivery. Further in vivo studies leading to the understanding of the mechanism of the 2'-OMe-4'-thioribonucleoside–modified AMO function as well as their potential side effect are currently under way.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

ACKNOWLEDGEMENT

The authors would like to thank Ms Y. Misawa (Hokkaido University) for technical assistance.

FUNDING

Grant-in-Aid for Scientific Research [23249008]; Grant-in-Aid for Scientific Research on Innovative Area “Nanomedicine Molecular Science” [2306] from the Ministry of Education, Culture, Sports, Science and Technology in Japan. Funding for open access: Grant-in-Aid for Scientific Research [23249008].

Conflict of interest statement. None declared.

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