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Published in:
Molecular Therapy - Nucleic Acids

Link to article, DOI:
10.1016/j.omtn.2019.05.024

Publication date:
2019

Document Version
Publisher’s PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):
Ray, R. M., Hansen, A. H., Slott, S., Taskova, M., Astakhova, K., & Morris, K. V. (2019). Control of LDL Uptake in Human Cells by Targeting the LDLR Regulatory Long Non-coding RNA BM450697. Molecular Therapy - Nucleic Acids, 17, 264-276. https://doi.org/10.1016/j.omtn.2019.05.024

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Control of LDL Uptake in Human Cells by Targeting the LDLR Regulatory Long Non-coding RNA BM450697

Roslyn M. Ray,1 Anders Højgaard Hansen,2 Sofie Slott,2 Maria Taskova,2 Kira Astakhova,2 and Kevin V. Morris1

INTRODUCTION

Hypercholesterolemia is a condition that is characterized by very high levels of cholesterol in the blood and is a major correlating factor with heart disease. Indeed, high levels of the low-density lipoprotein (LDL) have been causally linked to the development of atherosclerotic cardiovascular disease (ASCVD). A method to specifically reduce cholesterol in the blood in a long-term, stable manner could prove therapeutically relevant. Cholesterol is removed from the blood by the LDL receptor (LDLR) in the liver. Others and we have discovered that a long non-coding RNA (IncRNA; BM450697) functions as an endogenous epigenetic regulator of LDLR and that the repression of this IncRNA by the action of small interfering RNAs (siRNAs) results in the activation of LDLR. We found here, through the interrogation of two siRNAs that can target this IncRNA, both in a transcriptional and post-transcriptional manner, that BM450697 functions as a local scaffold for modulating LDLR transcription. Moreover, we found that conjugation of α-N-acetylgalactosamine (GalNAc) with two IncRNA-directed siRNAs allows for direct liver cell targeting of this IncRNA and functional enhanced uptake of cholesterol. Collectively, these data suggest that targeting the BM450697 IncRNA regulator of LDLR may result in a more specific, long-term, targeted approach to regulating cholesterol in the blood.

Hypercholesterolemia is a condition characterized by high levels of cholesterol in the blood and is a major factor that correlates with heart disease. Specifically, apolipoprotein-B (ApoB)-containing lipoproteins have been established as the causative agents in the development of atherosclerotic cardiovascular disease (ASCVD). Low-density lipoprotein (LDL) is a member of the ApoB lipoproteins, and reducing LDL levels in the blood may lead to a decreased risk of developing ASCVD. LDL blood levels are regulated by the LDL receptor (LDLR) in the liver, where active cycling of cholesterol from the blood occurs. Recent findings have suggested that long non-coding RNAs (lncRNAs) play an increasingly important role in cholesterol and lipid homeostasis. lncRNAs are non-coding transcripts that are over 200 nt in length and have been shown to modulate transcription or translation in a myriad of ways. For example, lncRNAs have been shown to modulate transcription through the recruitment of chromatin remodelling factors or by acting as mRNA decoys to sponge away or disrupt transcription factors or transcriptional machinery from binding to promoter sites or as enhancer RNAs to modulate transcription. In addition, lncRNAs have been shown to act as miRNA sponges, modulate splicing events, or alter activation states of proteins that lead to different transcriptional outcomes.

The modulation of LDL cholesterol occurs via a multi-step pathway. One branch of this pathway controls the transcription and recycling of the LDLR, whereas the other branch modulates cholesterol biogenesis, and the cross-talk between the pathways maintains intracellular cholesterol levels. Several lncRNAs have been found to participate in the modulation of cholesterol. For example, the lncRNAs liver-expressed LXR-induced sequence (LeXis) and macrophage-expressed LXR-induced sequence (MeXis) have been found to be integral in cholesterol biogenesis through modulation of the liver X receptor (LXR) pathway in mouse hepatocytes and macrophages, respectively, and both may contribute to the development of atherosclerosis. Further, overexpression of LeXis, using an adeno-associated virus (AAV) vector in mice, was found to significantly decrease cholesterol levels, suggesting that lncRNAs may be useful candidates for cholesterol maintenance. Matsui et al. found that small RNAs direct transcriptional gene activation (TGA) of LDLRs, indicating that a lncRNA is actively involved in LDLR regulation. This lncRNA (expressed sequence tag [EST] BM450697) is notably antisense (AS) to the LDLR promoter and is discordantly regulated relative to the LDLR, suggesting that this RNA may be an endogenous transcript involved in the direct regulation of LDLR transcription. Further, Matsui et al. found that small non-coding RNAs (lncRNAs) play an increasingly important role in cholesterol and lipid homeostasis.
determined that this lncRNA may be a useful therapeutic target for cholesterol modulation.

To this end, we sought to determine the molecular functioning of the lncRNA BM450697 to modulate LDLR mRNA levels and to identify candidate transcriptional-gene-silencing (TGS)-mediated small interfering RNAs (siRNAs) that may be therapeutically useful in modulating LDL levels. In addition, we used a N-acetylgalactosamine (GalNAc)-conjugated siRNAs as a delivery tool to directly target the liver for this siRNA-mediated effect. To determine mechanistically the involvement of the lncRNA BM450697 in LDLR regulation, we both overexpressed and repressed BM450697 expression. Notably, we found that BM450697 is an active transcriptional modulator of LDLR and that two siRNAs were observed to repress BM450697 and induce LDLR expression.

RESULTS
BM450697 Is a lncRNA That Is Located in Both the Nuclear and Cytoplasmic Fractions and Is Enriched at the Promoter of the LDLR Gene in the Form of DNA:RNA Hybrids

BM450697 is an AS RNA that overlaps the promoter of the LDLR gene according to the University of California, Santa Cruz (UCSC), Genome Browser (Figure 1A). According to the Annolnc database, BM450697 appears to be conserved in primates, but poorly conserved in vertebrates. Previous studies have observed that AS lncRNAs are functional in controlling the transcription of some gene promoters. Recent studies with the LDLR gene observed an AS lncRNA BM450697 (Figure 1A) that may be functional in RNA directed activation of LDLR. To interrogate this notion further, we assessed both Hep 3B and Hep G2 cell lines for the expression of BM450697 by directional RT and qPCR (Figure S1A). Indeed, we observed that both Hep G2 and Hep 3B liver cells express BM450697 (Figure S1A), which is AS to the LDLR promoter (Figure 1A). In addition, we find using a standard curve method that BM450697 has a low level of expression (around 45 copies/ng) of RNA (Figure S1B) in Hep 3B cells. To further
characterize this transcript, we confirmed that BM450697 is mostly polyadenylated, using poly deoxythymine (dT) magnetic beads to separate the RNA species (Figure S1C). Cytoplasmic fractionation showed that BM450697 was localized in both the cytoplasmic and nuclear fractions, using the nuclear IncRNA nuclear paraspeckle assembly transcript 1 (NEAT1) and the predominantly cytoplasmic IncRNA highly upregulated in liver cancer 1 (HULC1) as positive controls for each fraction (Figure 1B).

We used the chromatin immunoprecipitation by RNA pulldown (ChIRP) method to identify the loci associated with BM450697 (Figure 1C). Using several primer sets that tile along the LDLR promoter, we observed that BM450697 was enriched near the 3' end of the LDLR promoter near the translational start site (primer set 4). Further, these hybrids were lost with either RNase H treatment or under knockdown conditions. Similarly, we found that exogenously added BM450697 increased enrichment of these hybrids at the 3' end of the LDLR promoter (Figure 1D). These observations and those of others strongly support the notion that BM450697 is involved in transcriptional control of LDLR.

**BM450697 Modulates LDLR mRNA Levels through Transcriptional Interference**

We sought to determine the mechanism by which BM450697 may regulate LDLR mRNA levels. Our ChIRP and DRIP assays suggested that BM450697 binds directly to the LDLR promoter and, based on our work and that of others, has a low level of expression. Thus, we postulated that BM450697 may act by interfering with transcription factor SREBP1 or RNA polymerase II (Pol II) association and/or binding at the LDLR promoter (Figure 2A). SREBP1 is one of the main transcription factors that drives LDLR gene expression in the cholesterol pathway; thus, disruption of either SREBP1 or Pol II could result in a loss of LDLR transcription. We therefore performed chromatin immunoprecipitation (ChIP) assays with SREBP1a and Pol II (using the antibody toward the catalytically active phosphorylated serine 5 Pol II). We observed that when exogenous BM450697 RNA was transfected into Hep 3B cells, Pol II association at the LDLR promoter was significantly less compared with the control (scrambled, or a control lambda RNA; Figure 2B). Similarly, it appeared as if SREBP1a enrichment at the SREBP1 binding site was less than the control. However, this observation was not significant. Taken together, these data suggest that BM450697 decreases LDLR mRNA levels mechanistically by blocking interactions with Pol II and possibly SREBP1a at the promoter of LDLR.

**Repression of BM450697 by Small RNAs Induces the LDLR**

Previous observations in studies of promoter regulatory AS IncRNAs suggest that they can be targeted and repressed by the action of siRNAs and AS oligonucleotides. Rather than focus on the TGA of LDLR at the promoter site, we designed 10 siRNAs that could result in the TGS of the IncRNA, which could lead to an increase in LDLR expression, using a previously designed algorithm that generates siRNAs that target transcriptionally important regions. We interrogated the ability to target BM450697 with small RNAs in the population of IncRNA (Figures 3A and S2). Relative to both scrambled controls and an siRNA control (CTRL, 24UU) from Matsui et al., we found that the siRNAs 5 and p5 appeared to induce LDLR expression (Figure 3B). Notably, the previously described LDLR activating siRNA (24UU, CTRL) and the siRNAs 5 and p5 all demonstrated activation of LDLR in both Hep G2 and Hep 3B cells (Figure 3B). Interestingly, p5 demonstrated the most reproducible repression of BM450697 in both Hep G2 and Hep 3B.
We observed a ~30% reduction in BM450697 expression levels compared with the scrambled control with both 5 and p5 siRNAs in Hep3B cells. In addition, p5 resulted in a significant decrease in BM450697 levels in Hep G2 cells (Figure 3C). Using intron-exon spanning primers, we observed that si5 and p5 increased nascent LDLR mRNA transcripts (Figure 3C). It has also been established that LDLR levels are regulated, in part, by a negative-feedback loop that increases PCSK9 levels. Thus, we wanted to determine whether knockdown of BM450697 increases PCSK9 mRNA levels. We found that siRNA 5 and p5 did not significantly increase PCSK9 mRNA levels (Figure S3B). Collectively, these data, juxtaposed with those presented in Figures 1 and 2 and observed by others, suggest that BM450697 is a transcriptional regulator of LDLR and can therefore serve as a convenient and potentially highly specific target to control LDLR expression via directed small RNAs.

siRNA p5 May Function to Transcriptionally Control BM450697 Expression

We next wanted to elucidate whether any of these siRNAs that target BM450697 and result in the increase of LDLR mRNA expression levels are functional, by initiating TGS of BM450697. Using the ZENBU browser (http://fantom.gsc.riken.jp/zenbu) and the data obtained from Matsui et al., we found that several of our target siRNAs were within potentially important transcriptional regions for BM450697 expression (Figure S2). Thus, we performed both trichostatin A (TSA) and 5-aza experiments after siRNA treatment. TSA inhibits multiple histone deacetylases (HDACs), and whereas 5-aza inhibits DNA methyltransferase activity, both histone and DNA-methylation-associated proteins have been thought to be involved in the TGS pathway. Interestingly, siRNA 5 and p5, were found to be resistant to TSA treatment (Figures 4A and 4B), whereas the siRNA-mediated effects on LDLR mRNA levels were lost when treated with 5-aza in Hep 3B cells (Figure 4C). To confirm the 5-aza findings, we performed a ChIP assay to determine whether there was an enrichment of the methylation marker H3K27me3 at the siRNA target site (Figure 4D and S5). We observed a significant increase in H3K27me3 at the p5 target site in the p5-transfected condition after 72 h, whereas no increase in H3K27me3 was observed at the LDLR promoter site (Figure S4). Further, when assessing the effects of 5-aza on BM450697 mRNA expression, both si-5 and p5 silencing of the lncRNA were abrogated in the presence of 5-aza (Figure 4E). Taken together, our data suggest that we may be able to direct TGS to the promoter region for BM450697 (Figure 4C), which controls LDLR expression and, ultimately, LDL uptake by the liver.

siRNA-GalNAc Conjugates Gal-5 and -p5 Functions to Transcriptionally Control BM450697 Expression and Subsequently LDL Uptake in Hep 3B Cells

All natural 5 and p5 siRNAs were converted into their corresponding therapeutic analogs by insertion of 2′-OMe (2′-OMe) on the sugar moiety and phosphorothioate (PS) linkages in the backbone. Furthermore, the sense (S) strands of each pair were modified with GalNAc on the 3′ end to yield the corresponding modified siRNA-GalNAc conjugates of 5 and p5, i.e., Gal-5 and Gal-p5 (Table 1), respectively. The unconjugated controls (CTRL-5 and CTRL-p5) contained a free alkyne on the 3′ end, as designated in Table 1. It has been well established in the literature that GalNAcs are readily recognized and taken up by the asialoglycoprotein receptor (ASGPR) in hepatocytes. Upon ligand binding, the ASGPRs are endocytosed, whereupon the receptor is recycled back to the surface of the cell, after which the therapeutic siRNA can exert its function through the RNA interference (RNAi) pathway or association with argonaute 1 (AGO1) complexes, in the cell. Upon ligand binding, the ASGPRs are endocytosed, whereupon the receptor is recycled back to the surface of the cell, after which the therapeutic siRNA can exert its function through the RNA interference (RNAi) pathway or association with argonaute 1 (AGO1) complexes, in the cell.26,31 Dosing of the siRNA conjugates showed that both
Gal-5 and Gal-p5 had dose-dependent effects on LDLR mRNA and BM450697 RNA expression, with 50 nM showing significant effects for BM450697 repression and LDLR activation (Figures 5A and 5B). In addition, Gal-p5 appeared to be more potent than Gal-5, showing increased LDLR mRNA activation at 15 nM (Figure 5A). However, significant repression of BM450697 was observed only with 50 nM treatment with Gal-p5 (Figures 5A and 5B). It may be that a small reduction in the IncRNA is sufficient to elicit observable effects on LDLR mRNA expression, because of the low expression levels of this IncRNA in the cell (Figure S1B). In addition, we found a significant (~1.4-fold) increase in LDLR protein levels, with 50 nM Gal-p5 in Hep 3B cells (Figure 5C), suggesting that this conjugate increases both LDLR mRNA and protein levels in Hep 3B cells. Upon validation of the siRNA-GalNAc conjugates, we next sought to determine whether these siRNA conjugates had any physiological effect on cholesterol uptake in the Hep 3B cell line. We observed that Gal-5 and Gal-p5 had significantly higher levels of uptake than their respective controls (unconjugated siRNAs of CTRL-5 and CTRL- p5; Figures 5D and S7). The increased LDL uptake (Figure 5D) in the Gal-5 and Gal-p5 conditions (1.3- and 1.4-fold, respectively) appear to mimic the observed increase in LDLR protein levels in Figure 5C, suggesting that the increase in uptake may be proportional to the increase in LDLR protein levels. We observed similar effects when transfecting Hep 3B cells with the natural siRNAs, 5 or p5, in Hep 3B cells (Figure S6). Furthermore, Gal-5 and -p5 had similar levels of LDL uptake compared to 1 μM Lovastatin (the positive control). Taken together, these data suggest that Gal-5 and Gal-p5 modulated LDL uptake in this cell line model.

siRNA-GalNAc Conjugate Gal-p5 Increases LDLR mRNA Expression in Primary Hepatocytes

Bolstered by our findings that Gal-p5 appeared to increase LDLR mRNA levels, decrease BM450697 RNA levels, and increase LDL uptake, we wanted to determine if some of these effects were evident in primary human hepatocytes. We observed that direct addition of the Gal-p5 conjugate lead to an ~2.9-fold increase in LDLR mRNA expression, more so than the increase observed with the transfected natural siRNA p5 (Figure 5E) after 48 h. However, further optimizations are needed to improve the transfection efficiency in primary hepatocytes, as increased cytotoxicity related to the lipofectamine-based reagent may have masked the results after 48 h. Although these observations were not statistically significant, the trend suggests that the increase with Gal-p5 was similar to that with the positive control, 1 μM Lovastatin (Figure 5E). These data suggest that Gal-p5 is amenable to uptake in primary human hepatocytes and may increase LDLR mRNA levels ex vivo.

DISCUSSION

Our understanding of gene regulation by RNAs has been rapidly expanding. IncRNAs represent a class of RNAs that exert gene modulatory effects in both the nucleus and cytoplasm. Nuclear effects include recruitment of chromatin remodelling factors to either repress or de-repress gene expression, working as enhancer RNAs, through transcriptional interference at the promoter site, or acting as a scaffold for regulatory proteins at target sites in the DNA. BM450697 is an IncRNA described by Matsui et al. It is a 1450 nt transcript that runs AS to the LDLR promoter. According to the coding potential calculator (CPC2), BM450697 has a coding potential of 0.16, classifying this RNA as non-coding.

Based on our ChIRP and DRIP findings, this IncRNA is associated with the promoter of the LDLR gene in the form of DNA:RNA hybrids. We also determined that BM450697 is most likely to work through transcriptional interference, by preventing either transcriptional factor binding or Pol II binding to the promoter. Several studies have observed the effects of cis-acting IncRNAs in mediating transcriptional effects through interference. Martinez et al. observed that a non-coding transcript, upstream of the dihydrofolate reductase
(DHFR) gene, interacts directly with the promoter region to form stable DNA:RNA triple-helix hybrids, displacing Pol II and the basal transcription initiation complex from the DHFR promoter. They also observed that this non-coding transcript directly interacts with the general transcription factor IIB, to mediate this repressive function on DHFR mRNA transcription. Martens et al.10 observed that the ncRNA SRG1 in *Saccharomyces cerevisiae* represses D-3-phosphoglycerate dehydrogenase 1 (SER3) through SRG1 transcription through the SER3 promoter, which leads to a displacement of transcriptional activators and Pol II binding to the SER3 gene. Interestingly, Latos et al.36 found that the lncRNA *Airn* causes silencing, not through its lncRNA product, but rather through the transcription of the lncRNA itself, which leads to the displacement of Pol II at the promoter of the *Igf2r* gene. Interestingly, Latos et al.36 found that the lncRNA *Airn* causes silencing, not through its lncRNA product, but rather through the transcription of the lncRNA itself, which leads to the displacement of Pol II at the promoter of the *Igf2r* gene. The data reported here demonstrated that exogenously added BM450697 RNA led to a significant decrease in Pol II at the LDLR promoter. In addition, our ChIRP results suggest that lncRNA BM45069 is associated directly with the 3' end of the LDLR promoter, and our DRIP assay confirmed the presence of an DNA:RNA hybrid. It thus appears that our mechanism is most likely to occur through an DNA:RNA triple helix or hybrid at the LDLR promoter that interferes with Pol II binding, rather than by transcriptional interference through IncRNA transcription, which would only occur in cis.

Having established and expanded upon the role of BM450697 in LDLR mRNA transcription, we next determined its suitability as a therapeutic target. Targeting LDLR expression may offer a treatment for type II familial hypercholesterolemia (FH)16 and ASCVD.1 Currently, statins represent the first-line strategy, targeting FH by increasing LDLR levels. However, 2%–12% of those with FH are unable to tolerate high-intensity statins because of the adverse effects associated with statin use.37,38 Thus, an approach to directly targeting LDLR mRNA transcription may offer a suitable alternative to those unable to use statin therapy. Matsui et al.24 described that repressing BM450697 leads to a concomitant increase in LDLR expression, which we also observed. We found that siRNAs 5 and p5 appeared to be the most robust targets for LDLR expression and BM450697 silencing, and both did not significantly increase the expression in PCSK9 mRNA levels. Further, we observed that p5 repressed BM450697 RNA through a mechanism that may involve methylation related to the loss of the p5 effect with 5'0aza on both LDLR and BM450697 mRNA levels and an increased recruitment of H3K27me3 to the genomic target site of p5. The finding that siRNAs can mediate TGS through heterochromatin modification has been shown in several studies in yeast39,40 and in mammalian cells, either through AGO1 recruitment41,42 or through DNA methyltransferase 3A (DNMT3A) association at the target methylation site.26,43 Taken together, the observations presented here suggest that p5 may be useful as an siRNA therapeutic that may mediate long-term silencing events of lncRNA BM450697 that results in stable activation of LDLR.

We converted siRNA 5 and p5 into therapeutically relevant constructs primed for liver targeting (Gal-5 and -p5), by using the liver-specific GalNAc ligand as a targeting appendage attached to

### Table 1. Design of Candidate siRNAs-GalNAc Conjugates and Unconjugated Controls

| Oligonucleotide | Sequence (5' → 3') | siRNA          |
|-----------------|--------------------|----------------|
| 5X-S            | 5'-CGCGGCGAGGAGCAAGCGGACUU-3'X | Gal-5 (5Gal-S:5-AS) |
| 5Gal-S          | 5'-CGCGGCGAGGAGCAAGCGGACUU-3'-GalNAc | CTRL-5 (5X-S:5-AS) |
| 5-AS            | 5'-GUCGCCCUUGCUCCUCGCGGCGGUU-3' |                        |
| p5X-S           | 5'-CCCUGUGGAGACUUGGGGACUU-3'X | Gal-p5 (p5Gal-S:p5-AS) |
| p5Gal-S         | 5'-CCCUGUGGAGACUUGGGGACUU-3'-GalNAc | CTRL-p5 (p5X-S:p5-AS) |
| p5-AS           | 5'-GUCGCCCAAGGUCUCCACAGGGGCUU-3' |

S, sense; AS, antisense.
Figure 5. Chemically Modified siRNA-GalNAc Conjugates Gal-5 and -p5 Significantly Increase LDLR mRNA Expression in Hep 3B and Primary Hepatocytes, Decrease the IncRNA BM450697, and Increase LDL uptake in Hep 3B Cells

Hep 3B cells were seeded at a concentration of 100,000 cells/well. siRNA conjugates targeted toward the IncRNA (Gal-5 and Gal-p5) or their corresponding unconjugated controls (CTRL-5 or CTRL-p5) were added to the cells at increasing concentrations (15, 25, and 50 nM). After 48 h, the cells were harvested for RNA and reverse transcribed with either random hexamers (A) or a strand-specific primer (B). (A) Relative LDLR mRNA expression levels and (B) relative BM450697 mRNA expression levels. Each siRNA-GalNAc conjugate was normalized to its respective unconjugated control with each concentration set to 1. Data are the mean ± SEM of three independent experiments, performed in triplicate. (C) An LDL ELISA was used to determine LDL protein levels with 50 nM treated with galNAc-siRNA. (A) A two-way ANOVA with a post hoc Tukey test was used; ***p < 0.001. (B) An unpaired Student’s t test (*p < 0.05, **p < 0.01) and (C) a non-parametric one-way ANOVA with a post hoc rank test (*p < 0.05) were performed. (D) Primary hepatocytes were seeded at a concentration of 375,000 cells/well. Gal-p5 or its unconjugated control CTRL-p5 were added directly to the cells. Natural siRNAs (scrambled or p5) were transfected with RNAiMAX, as a control. After 48 h, the cells were harvested for RNA and 50 ng RNA was used in a one-step quantitative real-time PCR reaction. Relative LDLR mRNA expression levels Gal-5 or p5 was normalized to their respective unconjugated (CTRL-p5) or scrambled (Scr) control set to 1. Histograms are representative of the mean ± SEM of three independent experiments, performed in duplicate or triplicate.
therapeutic siRNA, Revusiran, in a phase I clinical trial and found that this subcutaneously administered conjugate was well tolerated and reduced expression of its target transthyretin (TTR) by approximately 90%, with a multiple-dose strategy. The researchers found that the conjugate was well tolerated in humans and demonstrated robust and efficient knockdown of its target gene in vivo. However, Fitzgerald et al.47 reported several mild-to-moderate side effects in 11%–15% of the population treated with the PCSK9 inhibitor. Our strategy to target a lncRNA with a discreet expression profile and effect, may offer an alternative strategy to reduce off-target effects that may result from targeting coding genes. However, this remains to be determined. Future studies would need to include a humanized-liver mouse or primate model to validate our findings in an in vivo setting, as BM450697 appears, like other lncRNAs,29 to be conserved to humans and primates.25 In addition, the use of humanized mouse models has been shown to better recapitulate the effects of human hepatocyte function.49,50 Further, we would need to validate the efficacy, LDL uptake and dosing strategy, and toxicity of our siRNA-GalNAc conjugate (Gal-p5) compared to other commercially available cholesterol-lowering strategies in different hepatic cell lines, as well as in primary human hepatocytes, to determine its true value as a therapeutic agent.

The results of our study suggest that targeting of lncRNAs may result in positive therapeutic outcomes. Indeed, several reviews26,44,45 and studies in atherosclerosis and cholesterol maintenance12–14 have suggested that lncRNAs are viable targets that have robust and pronounced effects when targeted through RNAi-mediated strategies.21–23,51–53 Invariably, there will be challenges in the delivery of RNAi-based therapeutics. However, the advent of the GalNAc-conjugated siRNA has shown great promise in liver-targeted diseases, with several GalNAc conjugates in phase I and II clinical trials.31 Our approach in targeting the TGS of BM450697 could be a promising strategy for long-term treatment of FH and those with hypercholesterolemia, where fewer doses of the siRNA-GalNAc conjugate are needed to maintain LDL levels in vivo.

MATERIAL AND METHODS

Cell Lines and Primary Hepatocytes

The human hepatocellular carcinoma cell lines Hep G2 and Hep 3B (ATCC, Manassas, VA, USA) were cultured in Eagle’s minimal essential medium (ATCC) supplemented with 10% fetal bovine serum (GeminiBio, Sacramento, CA, USA) at 37°C in a water-jacket incubator. The cells were mycoplasma free and were...
passaged twice weekly. Prepared and plateable primary human hepatocytes were purchased from Zen-Bio (Morrisville, NC, USA) and processed according to the manufacturer’s instructions, using human hepatocyte plating medium and human hepatocyte maintenance medium from the manufacturer (Zen-Bio). Primary hepatocytes were plated onto collagen-I-treated, 24-well plates purchased from Corning (Corning, NY, USA). The primary hepatocytes maintained their morphology for the duration of the study.

Antibodies, Plasmids, and All-Natural siRNAs
Pol II phosphorylated serine 5 C-terminal repeat domain (CTD; ab5131) and its respective rabbit IgG control antibody (ab171870), as well as mouse IgG controls (ab18413), were purchased from Abcam (Cambridge, UK). SREBP1 chip grade antibody (sc-13551X) was purchased from Santa Cruz (Dallas, TX, USA). S9.6 (MABE1095) was purchased from Merck Millipore (Burlington, MA, USA). A BM450697 expression vector was synthesized by GeneWiz (South Plainfield, NJ, USA) in a pcDNA3.1+ backbone. Ten all-natural siRNAs, designed to target the lncRNA SMB00392 or its putative promoter region, were synthesized by Integrated Technologies (Skokie, IL, USA). Based on initial screening analysis, two siRNA pairs (5 and p5) were chosen for further investigation.

Design of Modified siRNA-GalNAc Conjugates
Therapeutic siRNA Gal-5 and Gal-p5 (Table 1) were designed to contain alternating 2'-O-Me sugar modifications, along with PS linkages in the 3' ends. Contrary to S strands bearing only a single PS linkage at the GalNAc-modified 3' end, therapeutic AS strands were modified with three PS linkages at the 3' end, to improve resistance toward 3' exonuclease activity. Furthermore, AS strands were flanked at the 5' ends with two 3'-OMe modifications to improve stability toward 5' exonuclease, including alternating 2'-O-Me motifs for protection against endonucleases.49

Synthesis of 2'-OH/2'-OMe PO/PS siRNAs
The corresponding base- and backbone-modified versions of siRNA 5 and p5 were designed to also contain a serinol alkyne modifier on the 3' end of the S strands. Both pairs were synthesized in an N-dimethyltryptamine (DMT)-off mode on the Expedite 8909 (Perspective Biosystems, Framingham, MA, USA). RNA phosphoramide reagents were purchased from Glen Research (Sterling, VA, USA); other reagents for the oligonucleotide synthesis were obtained from ABI Scientific (Sterling, VA, USA). The synthesis was carried out in a 1 μmol scale using 3'-alkyne-modifier serinol CPG and NittoPhase UnyLinker Solid Support.

After synthesis, the column was dried and emptied into a PCR-clean vial (RNase and DNase free), together with filters, and incubated with 1 mL methylamine (33% in ethanol) at 65°C for 2 h. The mixture was allowed to reach room temperature and kept at 4°C for an additional 30 min. The mixture was filtered, evaporated, and treated with 100 μL desilylation reagent (750 μL N-methyl-2-pyrrolidone, 375 μL triethylamine, and 500 μL triethylamine trihydrofluoride) on a thermomixer at 67°C for 3 h. Then, 10 μL quenching reagent (10 mL of 3 M NaOAc and 10 μL diethyl dicarbonate) was added, followed by the addition of 15 μL of 5 M NaClO₄ and 10 μL of 3 M NaOAc. Cold acetone (1 mL) was added to precipitate the oligonucleotides. The vial was centrifuged at 1,000 rpm for 10 min and placed at 4°C for 30 min, and the supernatant was removed (washed three times). The pellet was dried on a speed-vac, redissolved in MilliQ, and subjected to ion exchange (IE) high-performance liquid chromatography (HPLC), using a Dionex UltiMate 3000 LC System equipped with a DNAPac PA-100 (9 × 250 mm; 0.5 M NaClO₄, 5%–60% over 45 min; 0.25 M Tris-Cl [pH 8], 10% over 45 min; MilliQ water). Collected fractions were combined and subjected to desalting column chromatography (Illustra NAP-5; GE Healthcare), and the purity of each oligonucleotide 5X-S, 5-AS, p5X-S, and p5-AS was confirmed by the analytical IE HPLC (>90% purity). The identity of oligonucleotides was confirmed by MALDI-TOF mass spectrometry (MS; Bruker MicroTOF-Q II mass spectrometer) in positive mode, using 3-hydroxypropionic acid matrix (10 mg/mL 3-hydroxypropionic acid and 50 mM ammonium citrate in 70% aqueous acetonitrile). MS-calculated/MS-found values ([M+H]+): 5Gal-S, 8210/8208; 5X-S, 7930/7928; p5X-S, 7831/7830. 5X-S, 7930/7928; p5X-S, 7831/7830.

Synthesis of siRNA-GalNAc Conjugate S Strands
All reagents for the click reactions were obtained from Sigma-Aldrich (St. Louis, MO, USA) and were used as received. All reagents were prepared as fresh solutions in MilliQ water prior to setting up click reactions. Conjugation with α-GalNAc-PEG3-azide (Sigma-Aldrich, SMB00392) was carried out in an inert atmosphere (argon). A solution of an alkyne-modified oligonucleotide 5X-S or p5X-S (20 nmol, 12 μL) was added to 142 μL MilliQ water, after which α-GalNAc-PEG3-azide (200 nmol, 7.6 μL), aminoguanidine hydrochloride (50 mM, 4 μL), N-trimethylammonium acetate (TEAA) buffer (1 M, pH 7.3, 20 μL), Cu(II) Tris((1-hydroxy-propyl-1H-1,2,3-triazol-4-yl)methyl)amine (THPTA; 10 mM, 10 μL), and acetic acid (50 mM, 4 μL) were added to get a total reaction volume of 200 μL. The reaction was stirred at room temperature for 12 h, after which the reaction mixture was subjected to gel filtration using a NAP-5 Sephadex column (Illustra; GE Healthcare, Pittsburgh, PA, USA). Purity was confirmed by the analytical IE HPLC at 20°C or 60°C above 90%). MS calculated/MS-found values ([M+H]+): 5Gal-S, 8308/8306; and p5Gal-S, 8210/8208.

ChIRP Assay
The ChIRP assay was performed as described by Chu et al.34,35 Briefly, 100 million cells were used per condition for the assay. The cells were harvested from 500 cm² dishes in 1× PBS and cross-linked with 0.5% methanol-free formaldehyde (Sigma-Aldrich) for 10 min, before quenching the reaction with 0.125 mM glycine for 5 min at room temperature. Cells were washed once with ice-cold 1× PBS (Ca and Mg free; Thermo Fisher Scientific, Waltham, MA, USA) and resuspended in 10 mL 1× radioimmunoprecipitation assay (RIPA) buffer (10 mM Tris-Cl [pH 8.0], 1 mM EDTA, 0.5 mM EGTA, 1% Triton X-100, 0.1% SDS, and 40 mM NaCl) with 1× Halt Protease Inhibitor.
Cocktail and 0.02 vol SUPERase In RNase Inhibitor (Invitrogen). The cells were resuspended with a 26G needle, and lysed at 4°C for 45 min on a rotator, and centrifuged at 10,000 x g for 10 min at 4°C. The supernatant was removed, and the nuclei were resuspended in 1 x micrococcal nuclease buffer (New England Biolabs). Two thousand units micrococcal nuclease (MNase) was added and the nuclei suspension was incubated at 37°C for 20 min. The MNase reaction was inhibited by using 0.1 vol 0.5 M EDTA. Nuclei were centrifuged at 10,000 x g for 10 min at 4°C and resuspended in 1 mL nuclear lysis buffer (1 x PBS, 1% SDS, 1% NP-40, and 1% Na-deoxycholate with 1 x Halt Protease Inhibitor and 0.02 vol SUPERase In RNase Inhibitor), resuspended with a 26G needle, and rotated for 30 min at 4°C. The nuclei were subsequently sonicated for 10 cycles (30 s on and 30 s off), to rupture the nuclei, and centrifuged at 10,000 x g for 10 min at 4°C, to remove the debris. The supernatant was collected, and 10% was collected as the input. The remainder of the nuclei lysate was used in the ChiRP assay.

Lysates were pre-cleared using Dynabeads MyOne Streptavidin C1 (Invitrogen) for 2 h at 4°C with rotation. Thereafter, biotinylated OS oligonucleotides (ASOs) toward BM450697 (Table S1) and scrambled biotinylated ASOs (Table S1) were used for each condition, to a final concentration of 1,200 pmol. Nuclei lysates were incubated in hybridization buffer (750 mM NaCl, 1% SDS, 50 mM Tris-Cl [pH 7.0], 1 mM EDTA, and 15% formamide) with 1 x Halt Protease Inhibitor Cocktail and 0.02 vol SUPERase In RNase Inhibitor. Hybridization occurred overnight at 37°C. Dynabeads MyOne Streptavidin C1 (Invitrogen) were washed according to the manufacturer’s instructions and resuspended in nuclear lysis buffer. Then, 200 µL of the prepared magnetic beads was used for each ChiRP condition. Magnetic beads were incubated with the lysates for 30 min at 37°C with rotation. The beads were collected on a magnetic stand and washed five times with 2 x saline sodium citrate (SSC) buffer (Invitrogen) with 0.5% SDS.

DNA was eluted off the beads, using 100 µg RNase A (Thermo Fisher Scientific) and 100 U RNase H (New England Biolabs) in 1 mL elution buffer (50 nM NaHCO3 and 1% SDS) at 37°C for 30 min with rotation. Thereafter, the elutes were collected, treated with proteinase K (New England Biolabs), and reverse cross-linked overnight at 65°C. DNA was isolated using the QIAGEN PCR purification kit.

The remainder of the reaction was used for RNA isolation with QIAzol (QIAGEN, Hilden, Germany), followed by a TURBO DNase treatment (Ambion, Invitrogen) for 30 min. Primer-specific RT was performed using the Quantitect Reverse Transcription kit (QIAGEN). qPCR was performed as previously described, using primers for the LDLR promoter (DNA) as well as primers specific for the lncRNA BM450697 (RNA; see Table S1 for the primer sets).

ChiP and DRIP Assay
Linearized BM450697 DNA or a lambda DNA control were transcribed using the Durascribe Synthesis Kit (Lucigen, Middleton, WI, USA). Thereafter, the DNA-free 2F RNA was dephosphorylated with calf intestinal phosphatase (New England Biolabs) for 30 min at 37°C. The RNA was subsequently purified with the RNA MEGAclear Kit (Ambion, Invitrogen), according to the manufacturer’s instructions.

For the ChiP assay, 2 million Hep 3B cells were seeded onto 15 cm² dishes. The next day, the cells were transfected with 10 nM BM450697 or the control lambda RNA or with 100 nM scrambled or p5 siRNA, and the media were replaced 24 h later. After 24 (RNA) or 48 (siRNA) h, the cells were cross-linked with 0.5% methanol-free formaldehyde for 10 min, before the reaction was quenched with 0.125 mM glycine for 5 min at room temperature. The cells were washed once with ice cold 1 x PBS and resuspended in 1 mL 1 x Halt Protease Inhibitor Cocktail. They were resuspended with a 26G needle, lysed at 4°C for 45 min on a rotator, and centrifuged at 10,000 x g for 10 min at 4°C. The supernatant was removed, and the nuclei were resuspended in 1 x micrococcal nuclease buffer (New England Biolabs). Four hundred units of micrococcal nuclease (MNase) was added, and the nuclei suspension was incubated at 37°C for 20 min. The MNase reaction was inhibited using 0.1 vol 0.5 M EDTA. The nuclei were centrifuged at 10,000 x g for 10 min at 4°C and resuspended in 1 mL nuclear lysis buffer (with 1 x Halt Protease Inhibitor Cocktail), resuspended with a 26G needle, and rotated for 30 min at 4°C. The nuclei were subsequently sonicated for five cycles (30 s on and 30 s off), to rupture the nuclei, and centrifuged at 10,000 x g for 10 min at 4°C, to remove debris. The supernatant was collected, and 10% was collected as the input. MNase-digested lysates were incubated with 2 µg antibody overnight with rotation at 4°C. Thereafter, 20 µL magnetic protein A/G beads (Thermo Fisher Scientific) was added to the lysates for 2 h at 4°C with rotation. Using a magnetic rack, bound beads were washed four times with a high salt buffer (0.1% SDS, 1% Triton X-100, 2 mM EDTA, 20 mM Tris-Cl [pH 8.1], and 500 mM NaCl), one time with LiCl washing buffer (0.25 M LiCl, 1% NP-40, 1% sodium deoxycholate, 1 mM EDTA, and 10 mM Tris-Cl [pH 8.1]), and one time with 1 x Tris-EDTA (TE) buffer (Ambion, Invitrogen). Samples were subsequently eluted in 100 µL elution buffer (1% SDS and 0.1 M NaHCO3) for 20 min at 68°C. The elutes were subjected to sequential RNase and proteinase K digestion for 1 h at 37°C, before reverse cross-linking overnight at 65°C. DNA was isolated by using a PCR purification kit (QIAGEN). To analyze the ChiP data, qPCR was performed with a standard curve in each run. The calculated concentrations were used to determine the enrichment of the protein at the target DNA site, as a fraction of input.

DRIP assays were performed according to Halász et al.56 Briefly, after treatment with either exogenously added 2F control or 2F BM450697 or with 100 nM scrambled or siRNA p5, cells were harvested for DNA using the Maxwell RSC Cultured Cells DNA Kit (Promega, Madison, WI, USA). Thereafter, cells were sonicated for 10 cycles of 30 s on and 30 s off, to shear the DNA to approximately 500 bp fragments. Thereafter, 5 µg DNA was used per condition, with half of the samples treated with RNase H (New England Biolabs) overnight at 37°C. RNase H-treated samples were purified using the DNA Clean and Concentrator kit.
(Zymo Research, Irvine, CA, USA). DNA samples were incubated with the 59.6 antibody, which specifically recognizes DNA:RNA hybrids (Merck Millipore), overnight at 4°C in IP buffer (50 mM HEPES/KOH [pH 7.5], 0.14 M NaCl, 5 mM EDTA, 1% Triton X-100, and 0.1% Na-deoxycholate), after which pre-blocked magnetic protein A/G beads were added for 2 h at 4°C. Samples were washed successively with 1 mL low-salt buffer (50 mM HEPES/KOH [pH 7.5], 0.14 M NaCl, 5 mM EDTA [pH 8], 1% Triton X-100, and 0.1% Na-deoxycholate), 1 mL high-salt buffer (50 mM HEPES/KOH [pH 7.5], 0.5 M NaCl, 5 mM EDTA [pH 8], 1% Triton X-100, and 0.1% Na-deoxycholate), 1 mL LiCl washing buffer (10 mM Tris-HCl [pH 8] and 10 mM EDTA [pH 8]) at 4°C and two washes of 1 mL TE (100 mM Tris-HCl [pH 8] and 0.5% NP-40, 0.5% Na-deoxycholate, and 1 mM EDTA [pH 8]), and two washes of 1 mL TE (100 mM Tris-HCl [pH 8] and 10 mM EDTA, and 1% SDS) for 15 min at 65°C. After purification, the nucleic acids were cleaned with the DNA Clean and Concentrator kit (Zymo Research, Irvine, CA, USA) in 30 μL TE buffer. The recovered DNA was then analyzed by qPCR, as mentioned above.

**RNA, cdNA, and Quantitative Real-Time PCR**

Cells were transfected with siRNAs (1–5 and p1–p5) using Lipofectamine RNAiMAX (Invitrogen) in Opti-MEM (GIBCO, Invitrogen). Briefly, the diluted siRNAs in Opti-MEM were added to RNAiMAX in Opti-MEM and incubated at room temperature for 20 min. Hep 3B cells were plated at a density of 75,000 cells/24-well plate and transfected the next day with the duplex siRNAs, using 1.5 μL RNAiMAX per condition. Hep G2 cells were reverse transfected at a density of 75,000 cells/24-well plate with 50 nM siRNA, using 2.4 μL RNAiMAX per condition. The media were replaced the following day, and the cells were harvested 72 h after transfection. In testing the modified siRNA-GalNAc conjugates (CTRL-5, CTRL-p5, Gal-5, and Gal-p5), the conjugates were added directly to the wells of either Hep 3B cells or primary hepatocytes, in the absence of any transfection reagent. Cells were harvested 48 h after transfection. RNA isolation was complex by using the SimplyRNA kit for RNA isolation (Promega), according to the manufacturer’s instructions. cDNA was synthesized by reverse transcribing 250 ng total RNA using the Quantitect Reverse Transcription kit, which includes a genomic DNA-wiping step (QIAGEN). Quantitative real-time PCR was performed with 2x Fast Universal qPCR Mix (Kapa Biosystems, Wilmington, MA, USA), according to the manufacturer’s instructions, on a Roche LightCycler 96 real-time PCR system (Roche, Basel, Switzerland). Thermal cycling parameters started with 3 min at 95°C, followed by 40 cycles of 95°C for 3 s and 60°C for 30 s. Specificity of the PCR products was verified by melting-curve analysis (primers used in this study can be found in Table S1). For primary hepatocytes LDLR and HPRT1 were amplified, using the Luna Universal One-Step qRT-PCR kit (New England Biolabs). Briefly, 50 ng RNA was added to each amplification reaction, with the following thermal cycling parameters: 10 min at 55°C for RT and 1 min initial denaturation at 95°C, followed by 40 cycles of 95°C for 10 s and 60°C for 30 s.

All data were analyzed by the 2^(-ΔΔCt) method, calibrated to a reference gene, and normalized to the scrambled-control- or control-treated sample set to 1.

**Poly(dt) Enrichment and Depletion**

Ten million Hep 3B cells were used per experiment, and poly(dt) RNA was selected for use with the Oligo(dt)25 Dynabeads magnetic capture kit, according to the manufacturer’s instructions (Thermo Fisher Scientific). The negative fraction was used as the poly(dt)-depleted fraction. One microgram of RNA was converted to cDNA in either BM450697-specific RT primer or with random hexamers, using the Quantitect Reverse Transcription kit (QIAGEN). PCR was performed as described above for BM450697 amplification. The PCR products were separated on a 1.5% Tris-acetate-EDTA (TAE) agarose gel at 100 V for 40 min and visualized using a Bio-Rad EZ Imager (Bio-Rad, Hercules, CA, USA).

**Subcellular Fractionation**

Ten million Hep 3B cells were used per experiment, in duplicate, according to the fractionation protocol for RNA by Gagnon et al.57 DNase-treated, total isolated RNA was converted to cDNA using the Quantitect Reverse Transcription kit (QIAGEN) with random hexamers or an RT-specific primer. In order to assess the quality of the fractionation, qPCR was performed, amplifying known nuclear (NEAT1) or cytoplasmic (HULC1) IncRNAs. BM450697 was amplified as described above (see Table S1 for primers). Relative amounts of each transcript were determined with a standard curve with Hep 3B genomic DNA and described as nanograms per reaction. The PCR products were separated on a 1.5% Tris-acetate-EDTA (TAE) agarose gel at 100 V for 40 min with a Bio-Rad EZ Imager (Bio-Rad).

**TSA and 5’ Aza Experiments**

Hep 3B and Hep G2 cells were seeded at a density of 100,000 cells/12-well plate, and transfected as previously described. Trichostatin A (Merck Millipore) was added to Hep 3B cells at a final concentration of 40 nM. The cells were harvested 72 h later for RNA, as previously described. 5’ Aza (Sigma-Aldrich, St. Louis, MO) was added daily to a final concentration of 7.5 μM, and the cells were harvested at 72 h for total RNA isolation.

**LDL Uptake Assay**

Hep 3B cells were seeded at a density of 75,000 cells/24-well plate. The next day, modified conjugates (CTRL5, CTRL-p5, Gal-5, or Gal-p5) were added directly to each well, whereas unmodified siRNAs (scrambled, 5, or p5) were transfected using 1.5 μL RNAiMAX per condition. Media were replaced at 48 h with
serum-free EMEM. The LDL uptake assay was performed according to the manufacturer’s instructions, using BODIPY-LDL at 2.5 μg/mL LDL per well (Molecular Probes, Invitrogen) the next day. Images were acquired with a Nikon ECLPSE TS100 fluorescence light microscope. Images were acquired using Nikon NIS Elements, and data were analyzed, using ImageJ, version 1.50i (Wayne Rasband; NIH, Bethesda, MD, USA). Lovastatin (Insolution Lovastatin, sodium salt; CalBiochem, Merck Millipore) at 1 μM was used as a positive control, with the medium replaced and fresh Lovastatin added each day. Heparin at 100 μg/mL was used as the negative control to prevent LDL uptake and was added 30 min prior to the assay (Thermo Fisher Scientific).

**LDLR ELISA**

Hep 3B cells were seeded at a density of 75,000 cells/24-well plate. The next day, modified conjugates (CTRL5, CTRL-p5, Gal-5, or Gal-p5) were added directly to each well, and 72 h later, total protein was harvested using RIPA buffer. LDLR protein levels were determined with a human LDLR Quantikine ELISA kit (R&D Systems, Minneapolis, MN, USA), according to the manufacturer’s instructions. Absorbance was measured on a Promega GloMax Explorer microplate reader (Promega).

**Statistical Analysis**

All experiments were performed in triplicate, with at least two biological experiments performed per data set. Graphing and statistical analyses were performed using Prism for Windows, version 7.03 (GraphPad Software, La Jolla, CA, USA). For experiments with multiple comparisons, one- or two-way ANOVA (parametric or non-parametric) was performed, with a post hoc Tukey’s, Dunnett’s, or non-parametric Dunn’s test. Where only two groups were compared, the two-tailed unpaired Student’s t-test was performed. Details of the graphical representation, statistics used, and n values for each figure are described in each legend.

**SUPPLEMENTAL INFORMATION**

Supplemental Information can be found online at https://doi.org/10.1016/j.omtn.2019.05.024.

**AUTHOR CONTRIBUTIONS**

R.M.R. and K.V.M. conceived of and designed the experiments; R.M.R. conducted the experiments; A.H.H, S.S, M.T., and K.V. and K.V.M. wrote the main manuscript and prepared the tables and figures. All authors reviewed the manuscript.

**CONFLICTS OF INTEREST**

R.M.R. and K.V.M. have patent PCT/US2018/016727. The remaining authors declare no competing interests.

**ACKNOWLEDGMENTS**

This project was supported by NIH NIAID grant P01 AI099783-01 and NIH grant DK104681-02 to K.V.M. The content is solely the responsibility of the authors and does not necessarily represent the official views of the NIH. We would like to thank the mass spectrometry core at City of Hope.

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