Mechanism of action of hepatitis B virus S antigen transport-inhibiting oligonucleotide polymer, STOPs, molecules

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A functional cure of chronic hepatitis B requires eliminating the hepatitis B virus (HBV)-encoded surface antigen (HBsAg), which can suppress immune responses. STOPs are phosphorothioated single-stranded oligonucleotides containing novel chemistries that significantly reduce HBsAgs produced by HBV-infected liver cells. The STOPs molecule ALG-10000 functions inside cells to reduce the levels of multiple HBV-encoded molecules. However, it does not bind HBV molecules. An affinity resin coupled with ALG-10000 was found to bind several proteins from liver cells harboring replicating HBV.

INTRODUCTION

Despite the availability of a highly effective vaccine, over 296 million people worldwide were living with chronic hepatitis B (CHB) in 2019.1 Untreated CHB can result in a lifetime of illness. Up to 40% of the patients will progress to liver cirrhosis and severe diseases.2 Persistent HBV infection is the result of the patient’s immune system failing to clear the virus infection. HBV produces various molecules during infection that can suppress innate and adaptive immunity, including the HBV surface (S) antigen (HBsAg).3,4 To increase the challenge to cure, HBV produces low copy numbers of covalently closed circular DNAs (cccDNAs) in the nucleus of the infected hepatocytes and can integrated partial forms of its genome into human chromosomes. Current classes of approved therapeutics for CHB, recombinant interferons and nucleos(t)ide analogues, are highly effective in controlling virion production but have low cure rates either separately or in combination.5,6,9-11

The difficulty of eradicating HBV has led to the concept of functional cure, the elimination of virus production and the clearance of immunosuppressive HBV molecules, especially HBsAg.9 Some patients with prolonged HBsAg loss seroconverted and displayed improved clinical outcomes for years, even though HBV DNA persisted in their livers.7-10 Numerous direct-acting antivirals targeting HBV entry, capsid assembly, cccDNA, HBV RNA, and HBsAg secretion are in development, with a goal toward a functional cure.10,12 Nucleic acid polymers (NAPs), which are phosphorothioate-containing oligonucleotides, are a promising class of therapeutics for treating CHB. NAPs can prevent the establishment of duck hepatitis virus infection and the release of the duck HBsAg from infected cells.13-15 In clinical trials, a significant proportion of patients treated with NAPs had more than a 5-log10 IU/mL decrease of HBsAg in their plasma 5 weeks or more after treatment onset.16,17 Some of the patients exhibiting seroclearance produced antibodies to HBsAg.17

How NAPs function to inhibit HBV infection and reduce HBsAg is not clear. In cultured cells, NAPs do not appear to activate innate immune responses.18 NAPs could also inhibit the hepatitis delta virus, a subviral agent that co-infects hepatocytes with HBV and uses the HBsAg to form infectious particles.19 Notably, NAPs do not demonstrate efficacy in animal models of HBV infection and efficacy was limited to human CHB patients. We have developed a new class of NAPs called S-antigen transport-inhibiting oligonucleotide polymers (STOPs). STOPs can potently inhibit HBV infection and HBsAg production in vitro.20,21 Here, we...
describe the elucidation of the mechanism of action of a STOPS molecule in inhibiting HBV infection and reducing HBsAg levels.

RESULTS

STOPS molecule ALG-10000 can potently reduce extracellular HBsAg produced by HBV

ALG-10000 consists of 20 repeats of adenosine and 5-methyl-cytosine locked nucleic acid (LNA) nucleosides connected by phosphorothioate linkages (Figure 1A). In cultured HepG2 cells expressing the HBV entry receptor sodium taurocholate co-transporting polypeptide (NTCP), ALG-10000 reduced extracellular HBsAg encoded by HBV in a concentration-dependent manner (Figure 1B). During an infection, extracellular HBsAg is a combination of secreted HBsAg oligomers and the envelope proteins on hepatitis B virions. The effective concentration for a 50% reduction of extracellular HBsAg (EC50) was 4 nM and the cellular cytotoxicity at 50% (CC50) was greater than 1 μM. In HepG2.2.15 cells that possess an integrated, tandem copy of the HBV genome, ALG-10000 reduced extracellular HBsAg levels with an EC50 of 3 nM and a CC50 of over 100 nM, the highest concentration tested (Figure 1C). STOPS can reduce extracellular HBsAg levels in HepG2.2.15 cells at better than 20-fold lower concentrations than can NAPs.20

Properties of STOPS in reducing HBV infection

HepG2.2.15 cells were used as a model system to understand how STOPS affect HBV replication and virion production. Initial key findings included that transfection of ALG-10000 into cells is required to reduce HBsAg levels. Post transfection, ALG-10000 localizes primarily to the cell cytoplasm but does not extensively co-localize with HBsAg. Furthermore, ALG-10000 does not activate nucleic acid-sensing innate immune receptors. These results are detailed in Figure S1.

We seek to understand the kinetics of HBsAg reduction in HepG2.2.15 cells treated with STOPS. ALG-10000 was transfected into the cells and the medium collected every 12 h. The amount of HBsAg in the medium was quantified and normalized to that of mock-treated control cells. The inhibition of extracellular HBsAg by ALG-10000 occurred in a biphasic manner that reached maximal levels about 72 h after the introduction of ALG-10000 (Figure 2A). The biphasic reduction of HBsAg suggests that ALG-10000 has a complex mechanism to inhibit extracellular HBsAg.

The reduced extracellular HBsAg could be solely due to ALG-10000 inhibiting HBsAg transport. In this case, ALG-10000 will not reduce the intracellular HBsAg level and may even increase it due to intracellular accumulation. To examine this, cell-associated HBsAg levels were assessed by western blots. Intracellular HBsAg was reduced in cells transfected with ALG-10000, and the reduction was more pronounced after 72 h (Figures 2B and S2A). The reduction of intracellular HBsAg suggests that ALG-10000 acts on a process or processes that precede the transport and trafficking of the HBsAg into the extracellular media.

We examined whether STOPS would inhibit HBV molecules in addition to HBsAg. Secreted HBV E antigen (HBeAg) was reduced after transfection of ALG-10000 (Figure 2C). The EC50 for HBeAg reduction was 2 nM, but the greatest extent of HBeAg inhibition (~75%) was reproducibly lower than that of HBsAg (>90%). The HBV polymerase was reduced by ALG-10000 (Figure 2D). Intracellular HBV core levels were also reduced but at a more modest level compared
ALG-10000 binds host factors needed for HBsAg production

STOPS could exert their inhibitory effect on HBV infection by binding one or more host factors required to produce HBV molecules. To identify these host proteins, we generated an ALG-10000 affinity resin and incubated it with solubilized extracts from HepG2.2.15 cells. After several stringent washes, proteins that remained bound were digested with trypsin and subjected to liquid chromatography-tandem mass spectroscopy (LC-MS/MS) and assignment to proteins. All proteins assigned with high confidence are derived from the host cell (Figure 3A); no HBV proteins were detected.

To confirm that the proteins that bind to ALG-10000 have a role in HBV infection, small interfering RNAs (siRNAs) were used to knock down the host mRNA/protein and the amount of extracellular HBsAg produced by HepG2.2.15 cells was determined. Four of the siRNAs targeting host proteins potently reduced the amount of extracellular HBsAg, each with an EC50 of less than 2 nM (Figure 3B). The siRNAs demonstrated no apparent cytotoxicity, even at the highest concentrations tested (Figures 3B and S3). siRNAs that target other proteins pulled down by ALG-10000 either had no effect on HBsAg levels or reduced HBsAg by less than 25%.

The four siRNAs targeted GRP78/HSP5A (glucose-regulated protein 78; heat shock protein family A (Hsp70) member 5), RPLP2 (ribosomal protein lateral stalk subunit P2), SRSF1 (serine-arginine splicing factor 1), and HNRNPA2B1 (heterogeneous ribonucleoprotein A2B1). RPLP2 functions in translating select membrane proteins.22 GRP78 is a chaperone for membrane-associated proteins and a regulator of the unfolded protein response.23 SRSF1 and HNRNPA2B1 participate in RNA processing, although they also have roles in other nucleic acid-associated processes.22,23,25 All four proteins have previously been described to function in viral infection.26–30

RPLP2 forms a heterodimer with RPLP1 that is attached to the ribosome through RPLP0.26,27 Our proteomic analysis did not identify with those of the HBsAg and HBeAg (Figure 2E). Finally, total HBV RNA in ALG-10000-treated cells was found to be reduced by only up to 50% (Figure 2F). Notably, an antisense oligonucleotide ALG-20002, which targets a sequence downstream of the HBsAg open reading frame (ORF), reduced the total HBV RNA to less than 5% (Figure 2F). ALG-10000 thus reduced the amounts of multiple HBV proteins and RNA; however, the levels of reduction differed. These differences suggest that STOPS use a complex mechanism to inhibit HBV protein and nucleic acid production.

ALG-10000 inhibition of HBV replication

(A) Kinetics of the inhibition of extracellular HBsAg by ALG-10000 and REP 2139. Each of the data points represents the HBsAg amount present in the HepG2.2.15 cell culture medium over a 12-h period. The amount is normalized to the levels in the mock-transfected cells. ALG-10000 was transfected into cells at 8 nM and REP 2139 at 20 nM. (B) Western blot image showing that ALG-10000 reduces intracellular large S antigen in HepG2.2.15 cells. L.C. denotes a loading control. (C) Dose response for the inhibition of HBeAg production by HepG2.2.15 cells. HBeAg cell viability shown in green and cell viability in black. (D) Western blot image showing that ALG-10000 can inhibit HBV polymerase accumulation. ALG-20002 was transfected at 5 nM. REP 2139 was transfected at 40 nM. The LC is the cellular GAPDH protein. (E) ALG-10000 can inhibit the accumulation of the HBV core in a concentration-dependent manner. (F) ALG-10000 can modestly reduce the accumulation of total HBV RNA present in the cell. HBV RNAs present in the cell lysate 72 h after transfection were quantified by detection of the HBx sequence found in all HBV RNAs. ALG-20002 is an antisense oligonucleotide that directly targets HBV RNA sequences for degradation. In the graphs in panel A, C, and F, each error bar represents one standard deviation of uncertainty.

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However, an siRNA that knocks down RPLP1 effectively reduced HBsAg levels with an EC50 of less than 2 nM (Figure 3B). Furthermore, a recombinant RPLP1 protein binds to the ALG-10000 affinity resin, although less well than recombinant RPLP2 (Figure 3C). The knockdowns of RPLP1 and RPLP2 resulted in a similar inhibition of HBsAg, suggesting that these proteins function in HBsAg production. RPLP1 is, therefore, the fifth host factor that binds ALG-10000 and contributes to the normal levels of extracellular HBsAg.

We sought to confirm the functional relevance of the identified host proteins. Recombinant proteins GRP78 and HNRNPA2B1 were found to bind to the ALG-10000 affinity resin (Figure 3C). In addition, the amounts of SRSF1, GRP78, and HNRNPA2B1 were reduced in cells transfected with ALG-10000 for 76 h (Figures 3D and S4). The levels of RPLP1 and RPLP2 were less affected by ALG-10000 at this time point (Figure 3D). However, their abundances were more obviously reduced in cells transfected with ALG-10000 for more than 76 h (data not shown). ALG-10000 may render bound proteins more susceptible to degradation.

We have further validated a role for GRP78 in HBV HBsAg production. GRP78 can function with a co-chaperone DNAJB9. siRNA knockdown of DNAJB9 decreased HBsAg levels (Figure S5A). In addition, deoxynivalenol, an inhibitor of GRP78, was found to reduce HBsAg levels (Figures S5B and S5C). Altogether these results confirm that GRP78, RPLP1, RPLP2, SRSF1, and HNRNPA1B2 interact with ALG-10000 and function in extracellular HBsAg production.

**STOPS optimally reduce HBsAg by binding to multiple host factors**

STOPs could bind to multiple cellular factors through a combination of their sequence, length, and/or higher-order structures. Truncating STOPs to be less than 32 nt in length was shown to severely decrease HBsAg production inhibition.20 To examine whether STOPs molecules possess secondary or tertiary structures, we examined whether ALG-10000 exhibits the hyperchromicity effect, an increase in absorbance upon denaturation. ALG-10000 absorbance was only negligibly affected by temperature, suggesting that it lacks base-pairing interactions and exists as a largely unstructured polymer (Figure 4A). The AC repeats in STOPs and the phosphorothioate backbone are likely significant features recognized by cellular proteins.

Altering the STOPs sequence should affect the proteins that it binds. To test this, we made STOPs molecules that contain one or more motifs that are more specifically recognized by SRSF1 and HNRNPA2B1.
These two proteins recognize an octameric splice junction sequence (Figure 4B).24,32 As a control, a molecule consisting of five scrambled octameric sequences was also tested. In HepG2.2.15 cells, the STOPS with one or two octamers had comparable or slightly improved EC50s for HBsAg reduction, as did ALG-10000. However, increasing the number of octamers reduced the maximal level of extracellular HBsAg inhibition from about 90% to 40% (Figure 4C). The molecule containing five scrambled motifs failed to inhibit extracellular HBsAg altogether. These results suggest that maximal STOPS activity requires the AC motifs and that STOPS bind to multiple proteins, not just RNA splicing factors.

The host factors had different contributions to processes important for HBV infection

We seek to determine whether the host factors could account for the various STOPS activities. siRNAs targeting the host factors were transfected into HepG2.2.15 cells, and the amount of different HBV molecules examined. While the knockdown of all five host factors reduced extracellular HBsAg, the kinetics of HBsAg reduction differed. Knockdown of RPLP1 and RPLP2 caused a more rapid decrease of extracellular HBsAg, suggesting that newly produced RPLP1 and RPLP2 can better act on HBsAg production (Figure 4C). The molecule containing five scrambled motifs failed to inhibit extracellular HBsAg altogether. These results suggest that maximal STOPS activity requires the AC motifs and that STOPS bind to multiple proteins, not just RNA splicing factors.

Intracellular HBsAg was reduced by the knockdown of all five host factors in HepG2.2.15 cells (Figures 5B and S6). The knockdown of all five host factors also reduced HBV polymerase levels (Figure 5C). In this experiment, the reduction of the HBV polymerase by siRNA targeting GRP78 was more modest, probably due to the low reduction of GRP78 in some samples.

The host factors had different effects on the levels of HBV molecules. The knockdown of RPLP1 and RPLP2 resulted in a greater reduction in HBsAg levels than did knockdown of other host factors (Figures 5D and 5E). Conversely, knockdown of RPLP1 and RPLP2 did not affect the total HBV RNA, while knockdown of SRSF1 and HNRNPA2B1 reduced the total RNA levels to 50% relative to the mock-treated samples (Figure 5E). The modest decline in the total HBV RNAs is dependent on the concentration of the siRNA targeting HNRNPA2B1 (Figure S7). Altogether, these results show that, while all five host factors participate in HBsAg and HBV polymerase production, only some are involved in the abundance of HBeAg and HBV RNA. Importantly, the five host factors that bind ALG-10000 can account for the differential reduction of HBV molecules in HepG2.2.15 cells treated with ALG-10000 (Figure 2).

The figure shows the absorbance of ALG-10000 under increasing temperature. The absorbance under temperature 40°C was measured with 2 μM ALG-10000 in three distinct buffers: PBS (blue line), PBS amended with 1 M NaCl (red line), and PBS amended with 2.5 mM MgCl₂ (green line).
extracellular HBsAg and HBeAg levels (Figures 6C and S8). The knockdown of SRSF1 reduced HBsAg levels in HBV-infected PHHs but had a more modest effect on HBeAg (Figure 6C). Unexpectedly, the knockdown of HNRNPA2B1 did not reduce extracellular HBsAg and HBeAg made by the HBV-infected PHHs (Figure 6C). Western blot of the PHH cell lysates showed that HNRNPA2B1 was reduced by the siHNRNPA2B1, ruling out the possibility that the siRNA was not functional (Figure S8B).

Since STOPS sequester host factors, reducing the host factors before HBV infection should prophylactically decrease productive infection. To test this notion, PHHs were transfected with STOPS or siRNAs to knockdown the host factors prior to HBV infection (Figure 6D). PHHs transfected with ALG-10000 and ALG-20002 exhibited significantly reduced levels of extracellular HBsAg and HBeAg (Figure 6E). The knockdown of GRP78, RPLP1, RPLP2, and SRSF1 before HBV infection reduced extracellular HBsAg and HBeAg (Figure 6E). In dose responses, the EC50 for HBsAg reduction was less than 1 nM (Figure S8C). REP 2139, which exhibited a higher EC50 in HepG2.2.15 cells, also exhibited a higher EC50 in HBV-infected PHHs. Again, the knockdown of HNRNPA2B1 did not affect extracellular HBsAg or HBeAg levels (Figure 6E). These results show that GRP78, RPLP1, RPLP2, and SRSF1 are important for HBV infection of PHH, while HNRNPA2B1 is not.

GRP78, RPLP1, and RPLP2 are required for the proper translation and folding of HBsAg

GRP78’s involvement in HBsAg production suggests that STOPS affect protein folding and degradation. Ubiquitination and proteasome degradation are the major pathways to degrade mis-folded proteins. Therefore, we examined whether STOPS could affect the ubiquitination of the HBsAg. The lysates from HepG2.2.15 cells transfected with ALG-10000 or siRNA were immunoprecipitated with antibody to HBsAg, eluted from the antibody, then subjected
to western blot analysis. The samples probed with an antibody to ubiquitin yielded a smear indicative of polyubiquitination (Figure S9). HBsAg ubiquitination in cells treated with ALG-10000 was approximately 2-fold higher than that of mock-treated samples (Figure S9). Immunoprecipitated samples probed with antibody to HBsAg showed a ladder of bands with a small increase in molecular weight (Figure 7A). Notably, HBsAg was already modified even in mock-treated cells but the ubiquitinated HBsAg increased in cells transfected with ALG-10000 or siRPLP2. These results show that HBsAg is normally ubiquitinated and that STOPS or a deficiency of RPLP2 can increase ubiquitination.

To better quantify HBsAg ubiquitination, we captured HBsAg from cell lysates on plates coated with anti-HBsAg antibody, and then detected ubiquitin with an antibody coupled to a reporter enzyme. The signal, normalized to the amount of intracellular HBsAg, demonstrated that ubiquitinated HBsAg increased by about 2-fold in ALG-10000-treated cells (Figure 7B). Ubiquitinated HBsAg was also increased by the knockdown of RPLP1, RPLP2, and GRP78, but not by the knockdown of SRSF1 (Figure 7C).

Proteasome inhibitors were used to examine whether STOPS affect the abundance of HBsAg through proteasome degradation. In cells treated with ALG-10000, proteasome inhibitors bortezomib or oprozomib did not affect extracellular HBsAg but resulted in a about 2-fold increase in the intracellular HBsAg (Figure 7D). Knockdown of siRPLP1 in HepG2.2.15 cells treated with bortezomib also increased intracellular HBsAg by more than 2-fold (Figure 7D). Finally, when the amounts of ubiquitinated intracellular HBsAg were increased in cells knocked down for RPLP1, RPLP2, and GRP78 and treated with bortezomib, the amount of ubiquitinated HBsAg was increased (Figure 7E). The knockdown of SFSR1, which likely does not directly function in HBsAg translation, did not increase the amount of ubiquitinated HBsAg. Altogether, RPLP1, RPLP2, and GRP78 participate in HBsAg production, and a reduction in their level in the cell resulted in increased ubiquitination and proteolysis of the HBsAg.

RPLP1 and RPLP2 require HBV RNA sequence to affect HBsAg production

Not all virus require RPLP1 or RPLP2 for the production of their structural proteins. We speculate that HBV may specifically recruit
these proteins to modulate the translation of the HBsAg. To examine this possibility, the ORFs of the large and small HBsAg were cloned into the expression plasmid pCDNA3.1. In HEK 293 cells, both proteins were expressed. However, the knockdowns of the host factors did not reduce the amounts of small or large HBsAg (Figures 8A and S10A).

To determine whether additional HBV sequence(s) can enable the host factors to affect HBsAg production, a plasmid containing the 1.05X HBV genome, p1.05Gen_psiCHECK2, was tested. In 293 cells, HBsAg produced from this plasmid was sensitive to the knockdown of GRP78, RPLP1, and RPLP2 (Figure S10B). ALG-20002 that targets a sequence of the HBsAg ORF also reduced HBsAg production, as would be expected. Knockdown of SRSF1 did not affect HBsAg levels. These results suggest that additional sequences present in the HBV genome are required for GRP78, RPLP1 and RPLP2 to affect HBsAg production.

A series of constructs containing different HBV sequences cloned into expression plasmid pCDNA3.1 were made to map the sequence(s) required for host factor activity. HBsAg expressed from a sequence from the start of the polymerase ORF to the HBV 3′ terminus in plasmid pPol-Gen_cDNA3.1 was sensitive to knockdown by RPLP1 and RPLP2 (Figure 8). Cells transfected with ALG-20002 also reduced the extracellular HBsAg expressed from this plasmid, as would be expected. HBsAg expressed from plasmid pPol-SAgter_cDNA3.1.
which lacks the sequence 3' of the HBsAg ORF, was unaffected by the knockdown of RPLP1 and RPLP2 (Figure 8). Finally, HBsAg expressed from plasmid pSAg-PsiI_cDNA3.1, which contains the HBsAg ORF and the HBV 3' sequence containing only a portion of the HBV epsilon sequence, was affected by the knockdown of RPLP1 and RPLP2 (Figure 8). These results suggest that sequence 3' of the HBsAg ORF, but not containing the intact epsilon element, contributes to engaging RPLP1 and RPLP2.

The knockdown of GRP78 reduced the amount of HBsAg produced from plasmid p1.05GEN_psiCHECK2, while its knockdown increased the amount of HBsAg produced from constructs that lack the sequence 5' of the polymerase ORF (Figures 8 and S10). GRP78 thus has a more complex role in HBsAg production that could involve some of its regulatory activities. These experiments show that the inhibition of HBsAg expression by STOPS is specific to HBV because sequences flanking the HBsAg ORF are required to engage RPLP1, RPLP2, and GRP78.

DISCUSSION

STOPS are a new class of NAPs that have the potential to dramatically reduce HBsAg in patients with chronic hepatitis. STOPS have significantly improved efficacy, being at least 20-fold more potent in vitro than previously described NAPs that could significantly reduce HBsAg in patient serum.16,17 Here, we have elucidated the mechanism of action of a prototype STOPS molecule, ALG-10000. ALG-10000 sequesters several cellular proteins needed to produce HBV proteins, including the HBsAg. Inadequate amounts of these cellular proteins resulted in reduced levels of HBsAg and increased the amounts of mis-folded protein that was degraded by the proteasome.

The chemical features of the STOPS molecules allow them to reduce HBV infection. Phosphorothioated oligonucleotides have higher affinities for the bound proteins than their phosphodiester counterparts.35 High-affinity binding by STOPS could render the bound proteins in conformations susceptible to being degraded. Host factors bound by ALG-10000 are present in reduced amounts, probably due...
to increased degradation. Reducing the amounts of the host proteins for molecular synthesis takes time and may account for the slow kinetics for HBsAg reduction. The AC repeats of STOPs contributes to the recognition by RNA-binding host proteins. Although SRSF1 and HNRNPA2B1 recognize a sequence containing the splice junction motif, they could also preferentially bind RNAs containing CA dinucleotide motifs.\textsuperscript{36,37} Finally, the 40-nt length of STOPs molecules may allow contact with multiple proteins. ALG-10000 lacks inter- or intramolecular base-pairing interactions, perhaps due to the methylene linkage between the ribose $2'$ oxygen and $4'$ carbon. While oligonucleotides containing LNAs strengthen binding to nucleic acid cells, or improperly modified amounts of GRP78 could result in increased amounts of mis-folded produce, fold, and post-translationally modify HBsAg. Reduced the amounts of RPLP1/2 could alter the proper timing needed to contribute to their mechanism of action.

STOPs binding to multiple host proteins accounts for the infection of HBV and HBsAg production. SRSF1 and HNRNPA2B1 function in RNA processing and RNA metabolism. STOPs and the host factor knockdown will reduce the total amounts of HBV only by up to 2-fold, suggesting that SRSF1 and HNRNPA2B1 may only act on select HBV RNAs. Consistent with this is that knockdown of SRSF1 and HNRNPA2B1 dramatically reduced HBsAg levels in HepG2.2.15 cells but only minimally reduced the HBeAg. It should be noted that HNRNPA2B1 is not needed for all cells infected by HBV. HNRNPA2B1 is a member of a family of proteins that can functionally complement each other.\textsuperscript{38} HNRNPA2B1 also recognizes m6A-modified RNA, and the m6A modification could vary with cells.\textsuperscript{24}

The major effect of STOPs on HBV infection was at the level of protein production and degradation. Viral structural proteins are typically made at high levels and may have additional requirements to their production, such as specialized translational factors and/or protein chaperones.\textsuperscript{40} For dengue virus, RPLP1/2 decreases the time for translational pausing that could be evolved to coordinate polypeptide synthesis and modification/processing.\textsuperscript{22,41} A reduction in the amounts of RPLP1/2 could alter the proper timing needed to produce, fold, and post-translationally modify HBsAg. Reduced amounts of GRP78 could result in increased amounts of mis-folded or improperly modified HBsAg that are subject to degradation. RPLP1 and RPLP2 are also required for the normal amounts of HBeAg and HBV polymerase. HBeAg has a signal peptide and is secreted after proteolytic processing, and it seems likely that RPLP1/2 and GRP78 are needed for its production and secretion. The HBV polymerase, however, is not a transmembrane protein. We speculate that RPLP1/2 and GRP78 may not be restricted to acting on transmembrane proteins.

Viruses are the masters of co-opting cellular factors for their own advantage. For the translation and production of HBsAg, HBV possesses additional RNA sequences to engage RPLP1/2 and GRP78. To our knowledge, RPLP1/2 and GRP78 have not been described to specifically bind RNA. Martinez-Azorin et al.\textsuperscript{11} have previously shown that RPLP1 and RPLP2 act to translate foot-and-mouth disease virus proteins that are made by internal ribosomal entry, but they did not demonstrate that RPLP1 and RPLP2 bind RNA. We cannot rule out that an accessory protein that binds RNA may bridge the contact between the HBV RNA and host factors GRP78 and RPLP1/2.

Antiviral compounds that act on cellular proteins have the advantage of decreasing the chance of the emergence of viral resistant mutations. However, a concern for molecules that act on cellular targets will be that they will demonstrate toxicity. We have not observed that effective concentrations of STOPs will cause toxicity \textit{in vitro}. Approved antivirals that act on cellular targets include maraviroc and the nucleotide analogue cytarabine.\textsuperscript{42} Modified oligonucleotides that target host molecules have also been approved to treat liver diseases or to reduce proteins made by the liver.\textsuperscript{43,44} The technology to preferentially target oligonucleotides to the liver, the site for HBV infection, is relatively mature.\textsuperscript{45} STOPs inhibition of HBsAg and HBeAg production and acceleration of HBsAg degradation could contribute to the functional cure of CHB.

MATERIALS AND METHODS

Key reagents

Deoxyxynivalenol (catalog # D0156) and cyanogen bromide-activated sepharose (catalog # S9142-1) are from Sigma-Aldrich. Bortezomib (catalog # ab142123) is from Abcam, and oprozomib (catalog # S7049) is from SelleckChem.

siRNAs that targeted host proteins were from the OnTARGETplus series from Horizon Discovery: siGRP78 (catalog # L-1008198-01), siRPLP1 (catalog # L-011135-00), siRPLP2 (catalog # L-004314-01), siSRSF1 (catalog # L-018672-01), siHNRNPA2B1 (L-011690-01), siDNAJB9 (catalog # L-012815-00), and siDNAJB12 (catalog # L-020585-00).

The majority of primary antibodies and horseradish peroxidase-conjugated secondary antibodies were from Abcam: anti-GRP78 (catalog # ab191023), anti-RPLP1 (catalog # ab121190), anti-RPLP2 (catalog # ab154958), anti-HNRNPA2B1(catalog # ab6102), anti-β-actin (catalog # ab8227), anti-GAPDH (catalog # ab8245), goat anti-rabbit-HRP (catalog # ab6721), and goat anti-mouse-HRP (catalog # ab6789). Anti-SRSF1 is from Invitrogen (catalog # 32-4600). Anti-HBsAg is from Fitzgerald Industries (catalog # 20-HR20). Anti-HBV polymerase is from Santa Cruz Biotech (catalog # Sc-81590). Anti-ubiquitin-HRP P4D1 is from Cytoskeleton (catalog # 140495).

Chemical synthesis of STOPs

Oligonucleotides, including STOPs, were synthesized on an ABI394 DNA synthesizer at 1-μmol scale using universal linker solid support. The synthesized material was deprotected and cleaved from the solid support by heating at 55°C in an ammonium hydroxide solution overnight or 65°C in a 1:1 ammonium hydroxide/methyl amine solution for 90 min. The reaction mixtures were filtered and evaporated in a Savant SpeedVac. The residues were reconstituted in water and...
purified using an ion exchange high-pressure liquid chromatography (HPLC) column (Thermo Scientific DNA Pac PA-100, 9 \times 250 mm) in a gradient of 25\%–85\% buffer B amended to buffer A (buffer A was 20 mM sodium phosphate, 10\% acetonitrile in water, pH 8; buffer B was 20 mM sodium phosphate, 1.8 M sodium bromide, 10\% acetonitrile in water, pH 8). The purified material was dried, reconstituted in water, desalted using Sephadex-25 columns, and eluted with DNase/ RNase-free distilled water. The final samples were characterized by analyzing the sample mass and purity using liquid chromatography-mass spectrometry (LC-MS) and reversed-phase HPLC. The final solutions were quantified by measuring absorbance at 260 nm using NanoDrop 2000c.

**Thermodenaturation studies**

All samples were prepared at 2 \muM concentration in Corning phosphate-buffered saline (PBS; catalog # 21-040-CV Corning). Where present, PBS was amended to contain 1 M NaCl, or 1 M NaCl and 2.5 mM MgCl2. The data were collected using a Shimadzu UV2600 spectrophotometer connected to a TMSPC-8 unit and recorded at a rate of 1 \degree C/min, in 1 \degree C intervals, for both the heating (20 \degree C–95 \degree C) and cooling (95 \degree C–20 \degree C) runs. Three independent trials were conducted for each molecule.

**Cell cultures and manipulations**

HepG2.2.15 cells that contain integrated tandem HBV genome and constitutively express HBsAg were grown and manipulated in a DMEM/F12 (1:1) media (catalog # 2021-05, Corning) containing 10\% fetal bovine serum (FBS) (catalog # F8192, Sigma), 1 \times nonessential amino acids (catalog # 34319012, Corning), and 1\% penicillin and streptomycin (catalog # 30-002, Corning). All manipulations used cells grown on collagen-coated 96-well plates cultured at 37 \degree C with 5\% CO2.

HepG2-NTCP cells were maintained in DMEM/F12 (catalog # 10-092-CM, Corning) with 10\% FBS, 1\% penicillin and streptomycin, 2 mM glutamine, 1\% nonessential amino acids, and 1\% sodium pyruvate. The cells were seeded at 20,000 cells/well in a 96-well plate. After the cells attached overnight, they were infected with HBV with 200 genome equivalents of HBV AD38 (genotype D; strain AYW). The infected cells were transfected with STOPS using Lipofectamine RNAiMAX on days 5 and 8. The HBsAg was measured in the supernatant on day 11, and the remaining cells constitutively express HBsAg were grown and manipulated in a minimal essential medium (MEM) medium supplemented with 10\% FBS, 1 \times nonessential amino acids, and 1 \times penicillin/streptomycin. The cells were typically plated at 40,000 cells per well into a 96-well plate.

**Oligonucleotide and plasmid transfection**

Each oligonucleotide to be transfected was diluted in Opti-MEM (catalog # 31980-070, Gibco) with 0.3 \muL of Lipofectamine RNAiMax (catalog # 13778-500, Invitrogen). HepG2.2.15 cells were typically transfected while the cells were plated. HepG2-NTCP and PHHs were transfected after the cells were attached to the plates. Plasmids were directly transfected into cells at a ratio of 100 ng/well and 0.3 \muL of Lipofectamine 2000 (catalog # 11668-019, Invitrogen).

**Protein detection and quantification**

Extracellular HBV HBsAg and HBeAg were quantified using ELISA kits (catalog # DS187701 and catalog # 05187703, DiaSino). The amounts of signal for HBsAg and HBeAg were assessed by chemiluminescence. HBV core protein was quantified using a colorimetric ELISA assay (catalog # VPK-150, Cell Biolabs). Chemiluminescence and color change was quantified using the Envision 2102 multi-label plate reader.

Western blots used cells that were lysed with ice-cold cell lysis buffer (catalog # 87787, Pierce) containing protease inhibitors (catalog # 04-693-132-001, Roche). The lysates used to quantify the amount of ubiquitination were modified with deubiquitnase inhibitors (catalog # NEM90BB, Cytoskeleton) according to the manufacturer’s instructions. The lysate was then treated with universal nuclease (catalog # 88700, Pierce) for 10 min to decrease their viscosity. Denaturing Laemmli loading buffer containing β-mercaptoethanol to 2 mM was added to a final concentration of 1 \times, and the lysate was incubated for 5 min at 85\degree C. Molecules in the lysate were separated by electrophoresis in 4\%–12\% NuPAGE Bis-Tris gel (catalog # NP0336, Invitrogen), followed by transfer to membranes and incubation with the primary and secondary antibodies. The western blot blocking and washes were performed with PBS-Tween 20 (catalog # 28352, Thermo Fisher). Western blot signals were developed with chemiluminescence substrate solution (catalog # 34579, Thermo Fisher) and imaged with a ProteinSimple imager. Quantification of the bands from the acquired images was performed using ImageJ software.
The HBsAg ELISA assay was adapted to quantify the ubiquitins attached to HBsAg. Lysates from the HepG2.2.15 cells were prepared in the lysis buffer containing both protease inhibitors and deubiquitinase inhibitors, and 30 μL of the lysate was diluted in Opti-MEM to a 100-μL volume that was then incubated with gentle shaking for 1 h at room temperature on an HBsAg capture plate. The plate was then washed five times with 250 μL of a wash buffer and incubated with a solution of anti-ubiquitin antibody conjugated to horseradish peroxidase for 1 h at room temperature. Following another 1-h incubation with gentle shaking at room temperature, the plate was washed five times and incubated with chemiluminescence substrate. The signal was quantified with an Envision 2102 multi-label plate reader.

RNA quantification

HBV RNA quantification used the Quantigene assay kit (Thermo Fisher catalog # QS0009) according to the manufacturer’s instructions. Briefly, cells were washed with cold PBS once, then lysed with the Quantigene lysis buffer containing protease K (Thermo Fisher, catalog # Q50512) at 37°C for 30 min. Twenty microliters of the cell lysate was used for hybridization with HBV RNA probe which targets the Hepatitis B Protein X (HBX) coding region (Thermo Fisher, catalog # SF-10405) for signal normalization. After an overnight hybridization on the capture plates, excess probes were washed away and the hybridization signal was amplified by hybridization with pre-amplifier, amplifier probe, and labeling probe. The chemiluminescence readout was determined using the Envision 2102 multi-label plate reader.

Quantification of the knockdown of siRNAs used real-time qRT-PCR. Total RNA was extracted from cells using the QuickExtract RNA extraction kit (catalog # QER090150, Lucigen) according to the manufacturer’s instructions. The cells were washed with cold PBS and lysed with RNA extraction solution by vortexing for 1 min, then centrifuged. The lysate was incubated at 65°C for 2 min, then treated with DNase at 37°C for 15 min, followed by incubation with the stop solution at 65°C for 10 min. RNA was quantified by qRT-PCR using qScript XLT One-Step RT-qPCR ToughMix (Quantabio, catalog # Q5132-500) and GRP78, RPLP1, RPLP2, or 18S TaqMan Gene Expression assays (catalog # 4448490). Two microliters of cell lysate was added to 20 μL of the reaction mix. qRT-PCR was performed on a qTower platform (Analytic-Jena) at 50°C for 10 min for reverse transcription, then 95°C for 1 min. DNA amplification used 40 cycles at 95°C for 10 s followed by 60°C for 45 s.

Confocal microscopy

HepG2.2.15 cells were grown on poly-L-lysine-coated coverslips to 60% confluency. After transfection, the cells were cultured for 24 h and then were fixed with 4% paraformaldehyde for 15 min at room temperature. The cells were washed with PBS twice and mounted on glass slides with anti-fade mounting medium and DAPI (catalog # P36941, Life Technologies), then dried overnight in the dark. Micrographs were acquired with a Leica TCS SP5 confocal inverted-base microscope with a 63× oil objective. The images were processed with the Leica LAS AF and ImageJ software. Colocalization of fluorophores was quantified using an ImageJ plug-in tool JACoP. 46

Proteomic analysis

The ALG-10000 affinity resin was made using the standard protocol suggested by the manufacturer of the CNBr-activated Sepharose resin (Sigma). Unreacted groups on the resin were quenched with 0.1 M Tris-Cl. A control resin used Tris-Cl as the ligand. After extensive washing in high salt, the resin that reacted with ALG-10000 contained over 125-fold more nucleic acid than the control resin.

The ALG-10000 resin and Tris resin were incubated with extracts from HepG2.2.15 cells solubilized by Dounce homogenization in a Tris-buffered saline containing 0.05% Tween 20 and protease inhibitors (catalog # 11873580001, Roche) for 1 h. The resin was then washed four times with the extraction buffer that contained increasing salt concentrations up to 1 M followed by a final wash with Tris-buffered saline. The resin and associated proteins were treated with trypsin (catalog # V5280, Promega) overnight, followed by termination of the reaction with 10% trifluoroacetic acid. Peptides were then purified using C-18 reverse-phase Zip-Tips (catalog # ZTC185096, EMD Millipore) and subjected to LC-MS/MS. Peptide quantification was performed by summation of the peptide ion intensities after normalization to the signal from the bradykinin fragment. Peptides were resolved by a Bruker Autoflex III MALDI-ToF mass spectrometer (Agilent Technologies) in positive ion mode. Assigned peaks all corresponded to within 0.5 Da of their theoretical masses. Database searches used the program Mascot (Matrix Science), with searches directed against the NCBI’s human proteome and viral proteomes.

Expression of recombinant HBsAg

The sequence of the HBV genotype D (GeneBank: NC003977.2) was used for cloning. The sequence used was numbered from 1 to 3,182; however, it should be noted that the HBV genome is circular and terminally redundant, and that cloning of HBV sequences for protein expression may span more than one genome. For the small S, the sequence from nucleotide 157 to 837 was chemically synthesized with flanking NheI and NotI sites and cloned into the same restriction sites in pcDNA3.1. cDNA to express the large S antigen was from nucleotide 2,850 to 837, flanked by the NheI and NotI sites, and was cloned into pcDNA3.1. This sequence has the initiation codon of HBV with a duplicated terminal sequence was chemically synthesized from nucleotide 1,808 to 2,000 flanked with Sall and NotI restriction sites. This was cloned into the Xhol and NotI restriction sites of the plasmid psiCHECK2.

DNAs to generate pPol-Gen-cDNA3.1 and pSAg-Gen_cDNA were chemically synthesized with flanking NheI and ApaI restriction sites. The insert was digested with restriction enzymes and ligated to pcDNA3.1 that had been cleaved with NheI and ApaI. The DNA to generate pSAg-PsiI_cDNA3.1 was generated from a
chemically synthesized DNA that contained an NheI site 5' of the large HBsAg ORF to the 3' termini of the HBV genome. The DNA was cleaved with the NheI restriction enzyme and a PsiI restriction enzyme that cleaves at nt 1,919 of the HBV genome. The cleaved DNA was ligated to pcDNA3.1, which had been cleaved with restriction enzymes NheI and EcoRV. Plasmids for transfection were prepared using the EndoFree Plasmid Maxiprep kit (catalog # 12362; Qiagen).

HEK 293 cells were seeded in 96-well plates at 4 x 10^5 cells per well. After attachment, the cells were transfected with oligonucleotides (10 nM ALG-10000, 5 nM ALG-20002, and 10 nM siRNA) and, 48 h later, super-transfected with 100 ng/well of plasmids encoding the HBsAg. The cell culture medium was harvested at 36 h post plasmid transfection to quantify the HBsAg by ELISA. Cell lysates were also collected at that time to examine the abundance of the cellular proteins.

SUPPLEMENTAL INFORMATION
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AUTHOR CONTRIBUTIONS
C.C.K. was responsible for the majority of the results in the manuscript and was primarily responsible for the writing of the manuscript. J.A.S. and L.B. provided critical guidance, editing, and approval of the work in the manuscript. Y.N. contributed the results in Figures 1B and 1C and generated two of the constructs used in Figure 8. S.R. contributed to the results in Figures 2F, 3F, and S7. T.S.D.C. contributed Figure 4A and synthesized the molecules tested in Figure 4B. S.R. and T.S.D.C. contributed to the editing of the manuscript. L.B., R.K.P., J.H., D.B.S., and L.B. established the design, chemical synthesis of the oligonucleotides, and their analytical and functional analyses. D.B.S. also edited the manuscript. L.M.B. conceived the project and provided critical analyses and resources for the project.

DECLARATION OF INTERESTS
All authors declare no competing interests.

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