Experimental Drugs for the Treatment of Hepatitis D

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Abstract: Chronic hepatitis D virus infection is the most severe form of viral hepatitis. Antiviral treatment is urgently needed to prevent patients from developing end stage liver disease or hepatocellular carcinoma. Treatment options were limited to off-label use of pegylated interferon alfa until conditional approval of bulevirtide by the EMA (European Medicines Agency) in July 2020. However, several other antiviral compounds are currently investigated and represent promising agents for the treatment of chronic HDV infection.

Keywords: hepatitis delta, HBV/HDV coinfection, interferon, bulevirtide, lonafarnib

Introduction

Chronic Hepatitis D Virus (HDV) infection, also known as hepatitis delta, is the most severe form of viral hepatitis.¹ Approximately 5% of all HBsAg positive patients are co-infected with HDV leading to 10–12 million cases worldwide.² However, hyperendemic regions experience higher rates of co-infection and screening rates may vary in different areas of the world. Chronic HDV infection (CHD) is associated with a 3.8-fold higher risk of developing liver cirrhosis compared to patients infected with hepatitis B virus (HBV) alone and patients experience a 1.28-fold higher risk for the development of hepatocellular carcinoma (HCC).³,⁴ All in all, the 5-year mortality rate is increased in patients with HBV/HDV coinfection compared to HBV mono-infected patients.⁵,⁶

Due to its progressive course, antiviral therapy for the treatment of HDV infection is urgently needed. Pegylated interferon alfa (PEG-IFN-α) is approved for the treatment of HBV and is also effective against HDV. In the two large controlled prospective HIDIT studies, the response rate was 23–48% at the end of therapy.⁷,⁸ However, twenty-four weeks after the end of therapy only about 25% of patients were HDV RNA negative and in the further long term about 50% of these patients had a late HDV-RNA relapse.⁹ Despite its limited long-term virological response rates, antiviral treatment with PEG-IFN-α is associated with a lower risk of developing clinical complications and maintained virological response is even more strongly linked to a more favorable long-term course.¹⁰,¹¹ Nevertheless, IFN-based antiviral treatment is accompanied by severe side effects, treatment is limited to patients with compensated liver disease and the aspired maintained virological response is rare. Therefore, there is an urgent need for new therapeutic approaches and several substances are currently evaluated in clinical phase II and III trials. Recently, the entry-inhibitor bulevirtide has received conditional approval by EMA while results of the ongoing phase III study are still pending. The prenylation inhibitor lonafarnib has entered phase III clinical trials. In addition, nucleic acid...
polymers, RNA interference compounds and interferon lambda are other potential substances currently being tested. Importantly, treatment with nucleot(s)ide analogues (NA), which inhibit reverse transcription of HBV, have no direct effect on HDV replication. Nevertheless, NA are considered in terms of liver disease severity and HBV DNA levels in the therapy of patients with HBV/HDV co-infection. An overview of therapeutic approaches is depicted in Figure 1 and detailed information on the different substances is provided in the following.

**Parameters for Treatment Response in Hepatitis Delta**

The aim of antiviral treatment of viral hepatitis is the prevention of hepatic complications like liver cirrhosis, hepatic decompensation, development of HCC and liver-related death. Surrogate markers to predict improved clinical outcome and therefore defining treatment response are still a matter of debate for CHD. In HBV mono-infection long-term suppression of HBV DNA is associated with a reduced risk of developing liver-related complications. Similarly, suppression of HCV RNA replication in patients chronically infected with HCV leads to better overall survival. For CHD, declines in HDV RNA during IFN-α treatment have been associated with improved clinical outcome even if HDV RNA negativity could not be achieved. Therefore, a group of experts recently suggested an HDV RNA decline of ≥2 log at the end of treatment as a surrogate parameter for treatment efficacy in CHD. However, HDV RNA typically fluctuates over

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**Figure 1** Therapeutic approaches for the treatment of chronic HDV infection are depicted in relation to the viral life cycle. (1) HDV virions attachment to heparan sulfate proteoglycans and binding of L-HBsAg pre-S1 region to the HBV/HDV specific receptor, hNTCP. Viral particles enter the cell through endocytosis and the viral ribonucleoprotein (RNP) is released in the cytoplasm. (2) Viral RNP translocation to the nucleus. (3) HDAg mRNA transcription and replication of HDV RNA. (4) L-HDAg contains a prenylation site that is farnesylated by a cellular farnesyltransferase before being re-translocated to the nucleus. (5) Both forms of HDAg interact with the newly synthesized genomic RNA to form new viral (RNP) that are exported to the cytoplasm. (6) Viral RNPs interact with the cytosolic part of HBsAg at the endoplasmic reticulum surface inducing their envelopment. (7) HDV virions are secreted form the infected cell. The different steps targeted by antiviral treatments are depicted. HBV co-infection is represented by presence of HBV cccDNA and integrated HBV DNA. The figure was created with BioRender.com.

**Abbreviations:** cccDNA, covalently closed circular DNA; HBsAg, hepatitis B surface antigen; HDV, hepatitis D virus; hNTCP, human sodium taurocholate co-transporting polypeptide; L-HDAg, large hepatitis D antigen; NAP, nucleic acid polymers; RNAi, RNA interference compounds.
time which creates uncertainty regarding the finality of this endpoint. Several secondary end-points like early virological response, biochemical response (ALT normalization at the end of treatment/during follow up), histological evaluation and changes of HBsAg are additional parameters, which should be assessed during treatment. Ideally, the endpoint of finite antiviral treatment would be the loss of HBsAg, which corresponds to functional cure in chronic HBV infection and would also result in the inability for HDV assembly, release and transmission. To achieve HBsAg clearance, it is important to consider that both cccDNA and integrated HBV DNA may be the source of HBsAg (Figure 1) and therefore must be addressed by either antiviral or immunomodulatory therapies. Since the endpoint HBsAg clearance is still very rarely achieved, negative HDV RNA 24 weeks after stopping antiviral treatment has been suggested as the endpoint of finite treatment strategies.16

**Therapeutic Agents**

In Figure 1 different therapeutic approaches for the treatment of CHD are depicted. Beside interferon alfa, the by far longest known agent, new substances and their mode of actions are illustrated. Table 1 provides an overview of antiviral compounds currently investigated in clinical trials.

**Interferon Alfa**

Since the 1980, interferon alfa, and later pegylated IFN-α, has been used as an off-label treatment of CHD. In the two largest prospective clinical trials viral response rates range between 25 to 30% without any clear additional benefit of concomitant treatment with nucleotide analogues.7,8 Virological response rates are further reduced since late relapses occur in up to 50% of patients with negative HDV RNA at the end of treatment.9 However, an antiviral treatment course might be worthy since HDV RNA suppression or decline due to treatment with PEG-IFN-α is associated with improved clinical long-term outcome according to several published studies.10,11,17

Positive long-term effects have to be weighed up against side effects like flu-like symptoms, anemia and thrombocytopenia or psychiatric disturbances which frequently occur during treatment. In addition to that, treatment is contraindicated in patients with advanced chronic liver disease therefore excluding many CHD patients urgently in need of antiviral treatment.

Treatment duration in most of the studies was one year and extension to 96 weeks did not improve virological response rates in the majority of patients.8 Therefore, international guidelines recommend a treatment duration of 48 weeks for PEG-IFN-α.13 An extension of treatment duration can be discussed if patients show an HBsAg decline assuming HBsAg loss during prolonged treatment. However, HBsAg loss only occurred in <10% of the patients during the large clinical trials and therefore remains a rare event.7,8 HDV RNA negativity at treatment week 24 was associated with negative HDV RNA 24 weeks after the end of treatment and could be utilized as a parameter for treatment management.18

**Bulevirtide**

Binding of HBsAg to the sodium taurocholate co-transporting polypeptide (NTCP) is crucial for HBV/HDV entry into hepatocytes.19 Bulevirtide (former Myrcludex B) is a small peptide blocking the engagement of the HBsAg preS1 domain with NTCP. Therefore, de novo infection of former uninfected hepatocytes is prohibited and over time recolonization of the liver with uninfected cells could finally lead to elimination of HDV.

In the first phase IIa study, 24 patients were included and equally randomized to receive (I) bulevirtide 2 mg (daily) alone for 24 weeks followed by PEG-IFN-α 180 μg (weekly) for 48 weeks or (II) the combination of bulevirtide 2 mg (daily) and PEG-IFN-α 180 μg (weekly) for 24 weeks followed by PEG-IFN-α monotherapy (weekly) for 24 weeks or (III) PEG-IFN-α alone for 48 weeks. HDV RNA declined significantly in all treatment groups at week 24. Monotherapy with bulevirtide was associated with HDV RNA decline in all patients and two patients became HDV RNA negative at week 24. Interestingly, combination with PEG-IFN-α increased the number of HDV RNA negative patients at week 24 as five patients of treatment group II showed negative HDV RNA. Furthermore, quantitative HDV RNA decline was more pronounced in the combination group. None of the patients achieved the primary endpoint of >0.5 log reduction of HBsAg at treatment week 12.20

In the following dose-finding phase IIb study, 120 patients were assigned to either receive bulevirtide 2 mg, 5 mg or 10 mg in combination with tenofovir daily or tenofovir daily alone for 24 weeks.21 The primary endpoint of >2log HDV RNA decline at the end of treatment was achieved in 46, 47 and 77% of patients with escalating doses of bulevirtide as compared to 3% in the TDF
Table 1 New Drugs for Antiviral Treatment in Chronic HDV Infection

<table>
<thead>
<tr>
<th>Compound</th>
<th>Target</th>
<th>Phase of Development</th>
<th>Comments/Data</th>
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<tbody>
<tr>
<td>Bulevirtide (entry inhibitor)</td>
<td>NCTP (Sodium taurocholate co-transporting polypeptide) receptor antagonist</td>
<td>Phase III</td>
<td>• Subcutaneous application</td>
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<td></td>
<td></td>
<td>Conditional approval by EMA</td>
<td>• Overall, well tolerated</td>
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<td></td>
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<td>• Increase of bile acids, local side reactions</td>
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<td></td>
<td></td>
<td></td>
<td>• Monotherapy: Decline of serum HDV RNA and ALT normalization, no effect on HBsAg (^{21})</td>
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<td></td>
<td></td>
<td></td>
<td>• Combination (pegIFNa): stronger effect on HDV RNA and HBsAg (^{22})</td>
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<tr>
<td>Lonafarnib (prenylation inhibitor)</td>
<td>Farnesyltransferase inhibitor</td>
<td>Phase III</td>
<td>• Oral application</td>
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<td></td>
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<td>• Boosting with ritonavir reduces side effects (GI side effects) (^{27})</td>
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<td></td>
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<td>• Monotherapy: HDV RNA decline, no effect on HBsAg</td>
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<td></td>
<td>• Combination (pegIFNa): stronger effect on HDV RNA</td>
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<td></td>
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<td></td>
<td>• Viral and biochemical flares post-treatment are associated with HDV RNA and ALT response (^{28})</td>
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<td>Pegylated interferon-lambda</td>
<td>Immune modulation</td>
<td>Phase II</td>
<td>• Subcutaneous application</td>
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<td></td>
<td>Induction of ISG</td>
<td></td>
<td>• Less side effects than pegIFNa</td>
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<td></td>
<td></td>
<td>• ALT flares on treatment (^{15})</td>
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<tr>
<td>REP 2139 (nucleic acid polymers)</td>
<td>Amphipathic alpha-helices in Class I surface glycoproteins</td>
<td>Phase II</td>
<td>• Intravenous application</td>
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<td></td>
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<td></td>
<td>• ALT flares in combination with TDF and pegIFNa</td>
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<td></td>
<td>• Strong effect on HDV RNA and HBsAg (^{28})</td>
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<td></td>
<td></td>
<td></td>
<td>• So far only data from 12 patients</td>
</tr>
<tr>
<td>JNJ-3989</td>
<td>Ribonucleic acid interference</td>
<td>Phase II</td>
<td>• Subcutaneous application</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>• Injection side irritations</td>
</tr>
<tr>
<td></td>
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<td></td>
<td>• No results regarding HDV treatment so far</td>
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monotherapy group (only one patient). More than 40% of patients treated with bulevirtide achieved ALT normalization but no dose depending effect could be detected as 43, 50 and 40% of the patients receiving bulevirtide 2 mg, 5 mg and 10 mg reached this secondary endpoint, respectively. An effect on HBsAg levels could not be revealed in any of the treatment groups and HDV RNA relapse after end of treatment occurred in most of the patients. Overall, treatment with bulevirtide was well tolerated: local reactions at the injection site and mild and transient neutropenia, thrombocytopenia and eosinophilia were observed. Elevation of bile salts did not lead to clinical symptoms.

In an additional multicenter phase IIb study prolonged treatment duration with bulevirtide at different dosages alone or in combination with PEG-IFN-α was assessed.\(^{22}\) 60 patients were randomly assigned to receive (I) PEG-IFN-α 180 µg, (II) bulevirtide 2 mg plus PEG-IFN-α 180 µg, (III) bulevirtide 5 mg plus PEG-IFN-α 180 µg or (IV) bulevirtide 2 mg for 48 weeks. Median HDV RNA reduction at the end of treatment was higher in the combination therapy groups as compared to PEG-IFN-α or bulevirtide 2 mg alone. Nevertheless, HDV RNA and ALT levels showed a linear decline in the bulevirtide monotherapy group during treatment. Similar to the previous phase II study, stopping of bulevirtide treatment caused HDV RNA relapse in >2/3 of the patients. Interestingly, 53.3% and 26.7% of the patients receiving bulevirtide 2 mg plus PEG-IFN-α or bulevirtide 5 mg plus PEG-IFN-α remained HDV RNA negative at 24 weeks after the end of treatment, respectively. HBsAg response defined as ≥1 log10 decline or negativity at 24 weeks after the end of treatment was detected in 46.7% of the patients treated with bulevirtide 2 mg plus PEG-IFN-α and in 20% receiving bulevirtide 5 mg plus PEG-IFN-α. None of the patients receiving bulevirtide or PEG-IFN-α monotherapy achieved this endpoint. In an extension study of this study patients received high dose bulevirtide (10 mg) either in combination with PEG-IFN-α or tenofovir. The primary end point of HDV RNA negativity 24 weeks after the end of treatment was achieved in 6.7% of the patients receiving bulevirtide 10 mg plus PEG-IFN-α and in 33.3% of the patients receiving the combination with tenofovir.\(^{23}\) None of the patients receiving bulevirtide plus tenofovir achieved HBsAg decline or negativity whereas one patient (6.7%)
of the becluviride 10 mg plus PEG-IFN-α group remained HBsAg negative 24 weeks after the end of treatment.

Overall, becluviride showed a dose-dependent antiviral effect regarding HDV RNA decline in the monotherapy trials. In combination with PEG-IFN-α the dose-dependent effect on HDV RNA and HBsAg decline reversed: The effect on HBsAg seemed to be more pronounced in the groups receiving lower doses of becluviride in combination with PEG-IFN-α. The reasons for this observation are not clear yet. However, becluviride and interferon show a synergistic antiviral effect in the treatment of CHD.

The results of the phase II studies, the limited therapeutic options for the treatment of CHD and its status of orphan disease finally led to the conditional approval of becluviride by the EMA (European Medical Agency) in July 2020. While phase III studies are still ongoing, patients with CHD, positive HDV RNA and compensated liver disease can now be treated with becluviride 2 mg as a monotherapy in the European Union. Duration of treatment is so far not defined. According to the EMA approval, treatment should be maintained as long as patients show a benefit. Some real-world experiences showed an ongoing biochemical and virological response in patients with long-term administration. However, data on prolonged treatment is still scarce.

**Lonafarnib**

Farnesylation of the large HD antigen is mandatory for HDV virion assembly. Inhibitors of cellular farnesylation disrupt viral assembly and therefore reduce the release of functional HDV virions. The farnesyl transferase inhibitor lonafarnib has been studied in phase I and II studies, and phase III studies are currently ongoing. Lonafarnib was tested as monotherapy, in combination with ritonavir (to increase lonafarnib levels by inhibition of cytochrome P450 3A4, the predominant mediator of lonafarnib metabolism) and with PEG-IFN-α. Results from the pilot study (LOWR HDV-1 study) showed that higher doses of lonafarnib were associated with a stronger decrease of HDV RNA. However, with increasing doses, adverse events, mainly gastrointestinal side effects like diarrhea and nausea, increased as well. The combination with ritonavir allowed lower lonafarnib doses, which resulted in better virological effects with lower rates of adverse events and a similar effect was observed in combination with PEG-IFN-α. The optimal treatment regimen was explored in the subsequent LOWR HDV-2 study. Fifty-five patients with compensated liver disease were included and randomized to either receive high (≥75 mg), medium (50 mg) or low (25 mg) doses of lonafarnib twice daily (bid) in combination with ritonavir or as a triple therapy in combination with PEG-IFN-α for 6 months. High rates of gastrointestinal side effects were observed in the high dose treatment arm. The primary end point of >2 log reduction of HDV RNA or negative HDV RNA at the end of treatment was achieved in 6 of 12 patients (50%) in the all-oral combination of lonafarnib 50 mg bid plus ritonavir treatment group while patients receiving 25 mg showed lower response rates. Triple therapy in combination with PEG-IFN-α achieved the best response rates and 8 of 9 patients showed a >2 log reduction of HDV RNA at the end of treatment. In a dose escalation study, the maximum dose of lonafarnib 100 mg bid plus ritonavir was tolerated by two third of the patients (10/15). At the end of the 24-week treatment, one patient showed negative HDV RNA and one patient had HDV RNA levels below the lower limit of detection. Liver enzymes were normalized in half of the patients. An additional study evaluated once-daily dosing of lonafarnib in combination with ritonavir. After 24 weeks of treatment with either 50, 75 or 100 mg lonafarnib boosted with ritonavir, six of 21 patients showed HDV RNA levels <250 IU/mL. Some patients showed post-treatment viral and biochemical flares, which were associated with future HDV RNA negativity, normalization of liver enzymes and in some cases suppression of HBV DNA. The mechanism of these responses after ending lonafarnib treatment is unclear but an immunological mechanism is suggested. The pronounced gastrointestinal side effects at high dosages, mainly nausea, diarrhea, anorexia and weight loss, are significantly reduced for dosages of 50 or 100 mg per day.

All in all, the available studies show that treatment with lonafarnib leads to HDV RNA decline and normalization of liver enzymes in patients with HDV infection. Boosting with ritonavir allows dose reduction of lonafarnib to reduce gastrointestinal side effects while maintaining antiviral efficacy. To date, the combination with PEG-IFN-α achieves the best response rates but results from larger clinical trials are still lacking. A phase III clinical trial has been initiated. In addition, lonafarnib is also evaluated in combination with pegylated interferon lambda (see below).
Pegylated Interferon Lambda

Pegylated interferon lambda (PEG-IFN-lambda) belongs to the type III interferons, which stimulate cell-mediated immune responses during viral infections. PEG-IFN-lambda binds to type III IFN receptors resulting in activation of the Jak-STAT signaling pathway, similar to IFN-alpha complexes. Type III interferon receptors are mainly expressed in hepatocytes and to a lesser extent in hematopoietic and central nervous system cells. Therefore, typical side effects of IFN-α treatment are not expected for treatment with PEG-IFN-lambda. PEG-IFN-lambda has already been studied in patients with chronic hepatitis B and C mono-infection. Treatment with PEG-IFN-lambda was associated with improved or similar rates of virologic response with fewer extrahepatic adverse events than IFN-α. However, on-treatment ALT and bilirubin flares have been observed. In a phase II study evaluating the safety, tolerability, and efficacy of PEG-IFN-lambda in CHD, 33 patients were randomized to receive either 120 µg or 180 µg subcutaneously for 48 weeks followed by a 24-week observation period. At the end of treatment, the primary end point of >2 log HDV RNA decline or negative HDV RNA occurred more frequently in patients treated with PEG-IFN-lambda 180 µg than 120 µg (7/14 vs. 4/19). Five of the 14 patients (36%) treated with PEG-IFN-lambda 180 µg remained HDV RNA negative after 24 weeks of follow-up. There were fewer adverse events than during treatment with PEG-IFN-α, but still some patients experienced hyperbilirubinemia, elevated liver enzymes and flu-like symptoms. Another trial investigating PEG-IFN-lambda in combination with lonafarnib over 24 weeks showed HDV RNA negativity or HDV RNA below the lower limit of quantification in 11 of 22 patients (50%) at the end of treatment. During follow-up relapse occurred in six patients leading to 23% of patients with maintained virological response 24 weeks after the end of treatment.

Nucleic Acid Polymers

Nucleic acid polymers (NAPs) inhibit the assembly and release of HBsAg coated viral particles and are therefore considered as an additional antiviral approach for the treatment of HDV infection. REP 2139 is the first NAP, which has been studied in clinical trials focusing on HDV treatment. In the first clinical study including 12 patients, 500 mg of REP 2139 was given intravenously once weekly over a period of 15 weeks followed by 15 weeks of 250 mg REP 2139 in combination with PEG-IFN-α followed by PEG-IFN-α monotherapy for 33 weeks. Eight patients showed HBsAg decline during REP 2139 monotherapy and 6 patients were HBsAg negative at the end of treatment. HDV RNA was significantly reduced during therapy and HDV RNA was negative in 75% of the patients (9/12) at the end of treatment. After a follow-up of 1 year, seven and five patients were HDV RNA and HBsAg negative, respectively. Five patients developed anti-HBs by the end of the 1-year follow up. Adverse events were reported by all patients, most of them experiencing neutropenia and thrombocytopenia as well as fever, chills and peripheral hyperemia. Furthermore, asymptomatic and transient elevation of liver enzymes have been reported. Studies investigating different dosing regimens are planned.

RNA Interference

RNA interference compounds (RNAi) are short RNA molecules targeting the transcripts of viral RNA. JNJ-3989 silences all transcripts deriving from HBV cccDNA and integrated viral DNA. Thus, siRNAs should lead to HBsAg decline. This could also impact HDV infection. So far, only HBV monoinfected patients have been included in clinical trials. In a phase IIA study, 40 patients were enrolled to monthly receive JNJ-3989 in different doses (subcutaneously) in combination with an oral nucleotide analogue for three months. HBsAg levels showed a rapid decline during treatment and 39/40 patients had a ≥1.0 log HBsAg reduction compared to baseline. 22 patients (22/39, 56%) showed sustained HBsAg reduction 9 months after the last RNAi dose. Adverse events were mostly injection site reaction. Overall, short-term treatment with the RNA interference therapy, JNJ-3989, resulted in sustained hepatitis B surface antigen suppression in patients with chronic hepatitis B receiving nucleotide analogue treatment. A phase II study to evaluate the safety and efficacy of JNJ-3989 in HBV infected patients who are co-infected with HDV is currently ongoing.

Conclusion

With the development of viral entry inhibitors, farnesylation inhibitors, viral particle formation blockers and interferon lambda, new promising treatment options are becoming available for patients with CHD. The entry-inhibitor bulevirtide received conditional approval by the
EMA this year and will change the current field of antiviral treatment of CHD. However, treatment duration is undetermined so far and prolonged treatment or even indefinite treatment are possible, especially in patients with advanced chronic liver disease. Since long-term treatment is currently investigated in the ongoing phase III study, results of this study are awaited eagerly. Data from phase II studies suggested a synergistic effect of the combination of buleviride with PEG-IFN-α but the optimal dose of buleviride for this concept has to be determined. Most importantly, maintained virological response was achievable with combination therapy which offers the potential for a finite antiviral treatment. Similar strategies are currently investigated for the other compounds either in combination with PEG-IFN-α or PEG-IFN-lambda.

The ultimate goal of CHD therapy is HBsAg clearance which also means functional cure of chronic HBV. So far, the highest rates of HBsAg loss were observed for nucleic acid polymers. To date, however, very few CHD patients have been treated with nucleic acid polymers and clearly more data is needed.

Importantly, all compounds have only been studied in patients with compensated liver disease. Since patients with CHD show high rates of advanced liver disease including a significant group of patients with decompensated liver cirrhosis, treatment options remain limited for these patients. Thus, further studies including this group of patients are necessary.

Finally, treatment strategies aiming to achieve HBsAg clearance need to be explored not only in chronic HBV but also in chronic HDV infection.

Disclosure
MC has received personal fees from AbbVie, Gilead Sciences, GSK, Janssen-Cilag, MSD Sharp & Dohme, Novartis, Spring Bank Pharmaceuticals, Swedish Orphan Biovitrum AB (SOBI). The authors report no other conflicts of interest in this work.

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