Original research

Randomised phase 2 study (JADE) of the HBV capsid assembly modulator JNJ-56136379 with or without a nucleos(t)ide analogue in patients with chronic hepatitis B infection

Harry L A Janssen,1,2 Jinlin Hou,3 Tarik Asselah,4,5 Henry L Y Chan,5 Fabien Zoulim,6 Yasuhito Tanaka,7 Ewa Janczewska,8 Ronald G Nahass,9 Stefan Bourgeois,10 Maria Buti,11 Pietro Lampertico,12,13 Oliver Lenz,14 Thierry Verbinnen,14 Joris Vandenbossche,14 Willem Talloen,14 Ronald Kalmeijer,15 Maria Beumont,15 Michael Biermer14, Umesh Shukla15

ABSTRACT

Objective We present the final analysis results of the phase 2 JADE study (ClinicalTrials.gov Identifier: NCT03361956).

Design 232 patients with chronic hepatitis B (CHB) not currently treated at study start (NCT) at study start or virologically suppressed were randomised to receive 75 mg (part 1) or 250 mg (part 2) JNJ-56136379, a hepatitis B virus (HBV)–capsid assembly modulator, one time per day or placebo with nucleos(t)ide analogue (NA) (tenofovir disoproxil fumarate/entecavir) or JNJ-56136379 alone (NCT-only) for ≥24 and ≤48 weeks.

Results In patients who are NCT hepatitis B e-antigen (HBeAg) positive, JNJ-56136379 75 mg+NA and 250 mg+NA showed limited mean (SE) hepatitis B surface antigen (HBsAg) declines (0.14 (0.10) and 0.41 (0.15), respectively) from baseline at Week 24 (primary endpoint): placebo+NA: 0.25 (0.11) log10 international unit (IU)/mL. In patients who are NCT HBeAg positive, mean (SE) HBV DNA declines at Week 24 were 5.53 (0.23) and 5.88 (0.34) for JNJ-56136379 75 mg+NA and 250 mg+NA, respectively, versus 5.21 (0.42) log 10 IU/mL (part 1) or 5.32 (0.33) log 10 IU/mL (part 2) JNJ-56136379, a class-variant nucleos(t)ide analogue in patients with CHB and was well tolerated.

Conclusions In patients with non-cirrhotic CHB, JNJ-56136379+NA showed pronounced reductions in HBV DNA and HBV RNA, limited HBsAg or HBeAg declines in patients who are NCT HBeAg positive, and was well tolerated, but no clear benefit with regards to efficacy of JNJ-56136379 over NA was observed.
with or without HBsAg seroconversion. Therapies include nucleos(t)ide analogues (NA), entecavir (ETV), tenofovir disoproxil fumarate (TDF) and tenofovir alafenamide or pegylated interferon alpha (peg-IFNα). These treatments lead to low rates of functional cure: ≈3% with NA and ≈10% with peg-IFNα. Treatment with peg-IFNα is associated with adverse effects, while NA therapy, which significantly reduces the risk of liver cirrhosis, decompensation and death, usually requires life-long treatment. There is a need for novel treatments for CHB that are of finite duration and increase functional cure rates to improve clinical outcomes, further reduce the risk of HCC, and address a significant stigma burden for patients.

One strategy is to combine HBV antiviral agents with different mechanisms of action (MoA) to intensify viral suppression and reduce levels of immune suppressive HBsAg. Capsid assembly modulators (CAMS) inhibit viral replication by interfering with HBV capsid assembly—a key step in virus production. CAM-N compounds, such as JNJ-56136379 and vecibcorvir (ABI-H0731), induce formation of morphologically intact but empty, non-functional capsids (N: normal structure). CAM-A agents, including RO7049389 (RG9709) and morphothiadin (GLS4)/ritonavir, result in the formation of pleiomorphic non-capsid structures (A: aberrant particle).

JNJ-56136379 binds to the HBV core protein and interferes with HBV viral replication at late and early stages via a dual MoA. The ‘primary’ MoA is interference with capsid assembly kinetics, preventing polymerase-pregenomic RNA (pgRNA) encapsidation and blocking HBV replication (median 50% effective concentration (EC50)/EC90 = 102 nM/376 nM). The ‘secondary’ MoA is inhibition of de novo covalently closed circular DNA (cccDNA) formation by interfering with disassembly of the capsid (median EC50/EC90 = 876 nM/4019 nM). NAs inhibit viral replication and virion production, but do not prevent encapsidation of pgRNA and release of HBV RNA-containing particles.

In a recently completed phase 1 study, JNJ-56136379 was well tolerated, with dose-proportional pharmacokinetics (PK) in healthy adults, and demonstrated antiviral activity with oral doses of JNJ-56136379 of 25–250 mg one time per day (qd) for 28 days in patients with treatment-naive CHB.

The final analysis results from the JADE study, evaluating efficacy, safety and PK of JNJ-56136379 administered alone or with an NA in patients with CHB are reported here.

**METHODS**

Additional methodology is included in the online supplemental methods.

**Study design and population**

JADE (NCT03361956) was a phase 2, randomised, partially-blinded, multicentre, interventional, placebo-controlled, two-part study in patients with CHB evaluating oral JNJ-56136379 administered as monotherapy or in combination with ETV or TDF (figure 1). Two, otherwise identical parts, explored 75 mg (part 1) and 250 mg (part 2) of JNJ-56136379 administered qd. The study included patients with non-cirrhosis aged 18–70 years with documented hepatitis B e-antigen (HBeAg) positive or negative CHB infection who were not currently treated at study start (NCT) or virologically suppressed (VS) with TDF or ETV at study start (inclusion and exclusion criteria provided in online supplemental table S1).

Each part comprised a screening (≤8 weeks), treatment (24 weeks plus a 24-week extension, depending on response) and follow-up phase (24 weeks or 48 weeks, depending on response). NCT patients were randomised 3:1:3 to receive open-label JNJ-56136379 monotherapy, placebo+NA or JNJ-56136379+NA. Patients who were VS were randomised 1:3 to receive placebo+NA or JNJ-56136379+NA. Randomisation was balanced and stratified according to HBeAg status (positive versus negative) and HBsAg level at screening (≥10 000 versus <10 000 international unit (IU)/mL for NCT patients who are HBeAg positive and ≥1000 versus <1000 IU/mL for all other patients).

Patients who completed the 24-week treatment phase and demonstrated a virological response by Week 20 (initially, HBV DNA <200 IU/mL; revised, HBV DNA <200 IU/mL), without any safety concerns, continued treatment for 24 weeks in a treatment extension phase. Patients meeting treatment completion criteria after 48 weeks were planned to stop all treatment and entered a 48-week follow-up phase. Patients undergoing 24 weeks of treatment not qualifying for treatment extension and patients who continued in the treatment extension phase and did not satisfy treatment completion criteria after 48 weeks, received NA monotherapy during a 24-week follow-up phase.

This study was conducted at 75 sites across 19 countries (Belgium, Canada, China, France, Germany, Hong Kong, Italy, Japan, South Korea, Malaysia, Poland, Russia, Spain, Taiwan, Thailand, Turkey, Ukraine, UK and USA). The study was conducted in accordance with the ethical principles outlined in the Declaration of Helsinki, International Conference on Harmonisation guidelines on Good Clinical Practices and applicable regulatory requirements, and approved by the relevant Independent Ethics Committee/Institutional Review Board. All patients provided written informed consent to participate in the study.

**Patient and public involvement**

Patients were invited by the investigator to participate in the study after ethics committee protocol approval. Patients were not involved in the study design or plans to disseminate study results. They were informed about the burden of intervention and could withdraw from the study at any time.

**Endpoints**

The primary endpoint was 1 log10 IU/mL mean decline in HBsAg levels from baseline to Week 24 for JNJ-56136379±NA versus placebo+NA. Secondary endpoints included antiviral activity on HBsAg, HBV DNA and HBeAg (in patients who are HBeAg positive only) measured during treatment (up to Week 48) and follow-up: (1) changes from baseline in levels, (2) proportion of patients with levels by predefined response category, and (3) proportion of patients with seroclearance and/or seroconversion.

**Study evaluations**

Study visits were scheduled at baseline and Weeks 1, 2, 4, 8, 12, 16, 20 and 24 during treatment and at Weeks 28, 32, 36, 44 and 48 during the treatment extension period. Patients who discontinued treatment prior to Week 48 or did not meet the treatment completion criteria at 48 weeks, had visits scheduled at follow-up Weeks 2, 4, 12 and 24 (end of study visit).
Levels of HBsAg, HBeAg, HBV DNA and HBV RNA were measured in plasma samples collected prior to dosing on Day 1 and at predefined time points throughout the study. Additional details for assessments are described in the online supplemental methods.

Safety
Safety and tolerability were assessed in all patients during study participation. Adverse events (AEs) and serious AEs (SAEs) were monitored. Physical examinations, vital signs measurements, 12-lead triplicate electrocardiograms and clinical laboratory tests were conducted at predefined time points.

Pharmacokinetics
Venous blood samples were collected to determine plasma concentrations of total JNJ-56136379 and NA using a validated liquid chromatography–mass spectrometry/mass spectrometry method. Urine samples were obtained at 24 hours in ~35% of the participants (PK subgroup at selected sites) to determine concentrations of JNJ-56136379 and NA.

HBV genome sequencing
HBV DNA was extracted from plasma samples, and next-generation sequencing (Illumina) with a 1% cut-off was performed for the full HBV genome.

Statistical analyses
No formal statistical hypotheses were evaluated. The target sample size was 220 patients, with 110 patients in each part (NCT: n=70; VS: n=40). Enrolment was planned to include ~40% of the NCT patients and ~30% of VS patients who were HBeAg positive. The probability to detect a treatment effect of 1 mean log\textsubscript{10} IU/mL decline in HBsAg from baseline to Week 24 for JNJ-56136379+NA versus placebo+NA, with ≥90% confidence, was 0.84 and 0.89 for patients who are HBeAg negative in the NCT and VS groups, respectively, and 0.64 and 0.57, respectively, for patients who are HBeAg positive.
Efficacy and safety analyses were conducted for all randomised patients who received ≥1 dose of study drug. Comparisons among the pooled placebo+NA, JNJ-56136379 75 mg+NA and JNJ-56136379 250 mg+NA treatment arms were described. Non-compartmental PK analyses of plasma and urine concentration-time data were performed. Descriptive statistics were used to compare plasma PK parameters for different treatments.

### RESULTS

#### Patient disposition and characteristics

The first patient was enrolled in February 2018; last observation for the last patient was in August 2020. Overall, 488 patients were screened and 232 (NCT: n=148; VS: n=84) were enrolled and randomised (figure 2). Patients were ineligible for the study due to failure to meet eligibility criteria (250 (51%)), patient withdrawal (4 (0.8%)), HBeAg enrolment target met (1 (0.2%)) and lost to follow-up (1 (0.2%)). Screen failure rate was mainly high in NCT patients (patients not meeting the NCT criteria). It was particularly high at the beginning of the study (≈88%) and decreased over time during study conduct. The placebo/ JNJ-56136379+NA treatment arms included 172 patients (NCT: n=88; VS: n=84), where 43, 66 and 63 patients were included in the pooled placebo+NA arms, JNJ-56136379 75 mg+NA and JNJ-56136379 250 mg+NA arms, respectively.

Sixty patients received JNJ-56136379 open-label monotherapy via staggered randomisation, where patients received 75 mg in part 1 and 250 mg in part 2 (75 mg: n=28; 250 mg: n=32). Following a viral breakthrough in five patients receiving the 75 mg dose between Weeks 16–20, JNJ-56136379 was discontinued in all 28 patients, and patients received NA. In the 250 mg monotherapy arm, viral breakthrough in one patient at Week 8 was followed by initiation of NA treatment plus ongoing JNJ-56136379 therapy in all patients.

Demographics and disease characteristics for patients in the NCT group of each treatment arm were broadly comparable, as were those for the VS patients in each arm (table 1 and online supplemental table S2). Across the treatment arms, the mean age was 39.7 years, 70% were men, 49% were Asian and 44% White and 37% were HBeAg positive. A numerically higher proportion of patients received TDF than ETV (69% vs 31%).

#### Figure 2 Patient disposition. The majority of patients were assessed with the initial extension criteria (<20 IU/mL). IU, international unit; NA, nucleos(t)ide analogue.
positive) were higher in NCT patients (2.33 (1.03) log10 IU/mL) than VS patients (0.39 (0.74) log10 IU/mL) in the combined JNJ-56136379 treatment arms. For NCT patients who were HBeAg positive, the mean (SD) baseline HBV DNA level was 7.86 (1.05) log10 IU/mL and 5.24 (1.32) log10 IU/mL for NCT patients who are HBeAg negative in the combined JNJ-56136379 treatment arms. Most VS patients (93% to 95%) had HBV DNA<lower limit of quantification (LLOQ) (20 IU/mL) and higher in NCT than VS patients (table 1).

Overall, 122/129 (95%) patients receiving JNJ-56136379+NA and 41/43 (95%) receiving placebo+NA completed Week 24 study treatment; 91/129 (71%) and 33/43 (77%), respectively, entered the 24-week treatment extension phase (figure 2), which included 47/88 (53%) NCT patients and 77/84 (92%) VS patients (online supplemental figure S1). Most patients who entered the 24-week treatment extension phase completed the extension (figure 2). No patients met the treatment completion criteria at Week 48.

### Antiviral efficacy

#### Primary endpoint

In NCT patients who are HBeAg positive, JNJ-56136379 75 mg and 250 mg+NA resulted in a mean (SE) HBsAg decline from baseline at Week 24 of 0.14 (0.10) and 0.41 (0.15) log10 IU/mL, respectively, versus 0.25 (0.11) log10 IU/mL for placebo+NA (table 2; figure 3A; figure 4; online supplemental figure S2). The proportion of NCT patients who are HBeAg positive with >0.3 log10 IU/mL reduction (corresponding to a 50% reduction) in HBsAg from baseline was 4/12 (33%) (JNJ-56136379 75 mg+NA), 5/11 (45%) (JNJ-56136379 250 mg+NA) and 2/8 (25%) (placebo+NA) (table 2). In NCT patients who are HBeAg negative or VS (HBeAg positive or negative) patients, there was no relevant effect of JNJ-56136379 75 mg and 250 mg+NA or placebo+NA on mean HBsAg levels (figure 3A).

#### Secondary exploratory endpoints during 24 weeks of combination treatment

**HBV DNA**

Change from baseline in HBV DNA levels was assessed in NCT patients (VS patients typically had levels <LLOQ at baseline). In NCT patients who are HBeAg positive (mean baseline HBV DNA levels, 7.65–8.24 log10 IU/mL), there were pronounced declines in mean (SE) HBV DNA at Week 24 of 5.53 (0.23) and 5.88 (0.34) log10 IU/mL for JNJ-56136379 75 mg and 250 mg+NA, respectively, and 5.21 (0.42) log10 IU/mL for placebo+NA (table 2; figure 5A; online supplemental figure S3A). In NCT patients who are HBeAg negative, interpretation of mean (SE) HBV DNA decline was confounded due to many patients (n=22/89 (25%) with HBV DNA levels <LLOQ from Week 4 onwards (online supplemental table S3).
### Table 2  Summary of antiviral efficacy at Week 24 (pooled placebo/JNJ-56136379+NA treatment arms)

<table>
<thead>
<tr>
<th>Population</th>
<th>NCT (n=22)</th>
<th>VS (n=21)</th>
<th>NCT (n=33)</th>
<th>VS (n=33)</th>
<th>NCT (n=33)</th>
<th>VS (n=30)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HBeAg status</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>+ (n=8)</td>
<td>− (n=14)</td>
<td>+ (n=6)</td>
<td>− (n=15)</td>
<td>+ (n=12)</td>
<td>− (n=21)</td>
<td>+ (n=9)</td>
</tr>
<tr>
<td>HBeAg, log10 IU/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Change from baseline at Week 24, mean (SE)</td>
<td>−0.25 (0.11)</td>
<td>0.02 (0.02)</td>
<td>0.01 (0.06)</td>
<td>0.02 (0.02)</td>
<td>−0.14 (0.10)</td>
<td>0.04 (0.02)</td>
</tr>
<tr>
<td>Patients with &gt;0.3 log10 decline, n (%)</td>
<td>2/8 (25)</td>
<td>0/13 (0)</td>
<td>0/5 (0)</td>
<td>0/15 (0)</td>
<td>4/12 (33)</td>
<td>0/21 (0)</td>
</tr>
<tr>
<td>Patients with &gt;0.5 log10 decline, n (%)</td>
<td>1/8 (13)</td>
<td>0/13 (0)</td>
<td>0/5 (0)</td>
<td>0/15 (0)</td>
<td>1/12 (8)</td>
<td>0/21 (0)</td>
</tr>
<tr>
<td>HBV DNA, log10 IU/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Change from baseline at Week 24, mean (SE)</td>
<td>−5.21 (0.42)</td>
<td>−3.62 (0.36)</td>
<td>−5.33 (0.06)</td>
<td>−0.03 (0.06)</td>
<td>−2.00 (0.52)</td>
<td>−1.41 (0.18)</td>
</tr>
<tr>
<td>Patients with HBV DNA &lt;LLOQ, n (%)</td>
<td>1/8 (13)</td>
<td>12/13 (92)</td>
<td>5/5 (100)</td>
<td>15/15 (100)</td>
<td>0/12</td>
<td>14/21 (67)</td>
</tr>
<tr>
<td>HBV RNA, log10 copies/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Change from baseline at Week 24, mean (SE)</td>
<td>−0.81 (0.28)</td>
<td>−0.31 (0.24)</td>
<td>−0.49 (0.09)</td>
<td>−0.32 (0.16)</td>
<td>Not applicable</td>
<td>−0.70 (0.20)</td>
</tr>
<tr>
<td>Patients with &gt;0.3 log10 decline, n (%)</td>
<td>5/8 (63)</td>
<td>Not applicable</td>
<td>1/5 (20)</td>
<td>Not applicable</td>
<td>10/12 (83)</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Patients with &gt;0.5 log10 decline, n (%)</td>
<td>5/8 (63)</td>
<td>Not applicable</td>
<td>1/5 (20)</td>
<td>Not applicable</td>
<td>5/12 (42)</td>
<td>Not applicable</td>
</tr>
<tr>
<td>Patients with &gt;1.0 log10 decline, n (%)</td>
<td>3/8 (38)</td>
<td>Not applicable</td>
<td>1/5 (20)</td>
<td>Not applicable</td>
<td>1/12 (8)</td>
<td>Not applicable</td>
</tr>
<tr>
<td>HBcrAg, log10 U/mL</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Change from baseline at Week 24, mean (SE)</td>
<td>−0.84 (0.27)</td>
<td>−0.91 (0.21)</td>
<td>−0.08 (0.04)</td>
<td>−0.28 (0.12)</td>
<td>−0.68 (0.12)</td>
<td>−0.70 (0.13)</td>
</tr>
<tr>
<td>Patients with HBcrAg &lt;LLOQ, n (%)</td>
<td>0/8</td>
<td>3/14 (21)</td>
<td>0/6</td>
<td>4/15 (27)</td>
<td>0/12</td>
<td>2/21 (10)</td>
</tr>
<tr>
<td>Patients with &gt;0.3 log10 decline, n (%)</td>
<td>5/8 (63)</td>
<td>9/13 (69)</td>
<td>0/5</td>
<td>4/15 (27)</td>
<td>10/12 (83)</td>
<td>14/21 (67)</td>
</tr>
<tr>
<td>Patients with &gt;0.5 log10 decline, n (%)</td>
<td>4/8 (50)</td>
<td>8/13 (62)</td>
<td>0/5</td>
<td>4/15 (27)</td>
<td>7/11 (63)</td>
<td>11/21 (52)</td>
</tr>
<tr>
<td>Patients with &gt;1.0 log10 decline, n (%)</td>
<td>0/8</td>
<td>8/13 (62)</td>
<td>0/5</td>
<td>7/11 (63)</td>
<td>3/12 (25)</td>
<td>4/11 (36)</td>
</tr>
</tbody>
</table>

*Seven patients in the placebo+NA group, 13 patients in the JNJ-56136379 75 mg+NA group and eight patients in the JNJ-56136379 250 mg+NA group had HBV RNA detectable at baseline, and 1/7 (14%), 13/13 (100%) and 8/8 (100%), respectively, were TND at Week 24. HBeAg values >ULOQ were set to 5.1 log10, HBV DNA values < LLOQ (20-100 IU/mL) target detected or target not detected were imputed respectively with 15 (IU/mL or 5 (IU/mL); not detected HBV RNA values were imputed by a value of 5 copies/mL; HBeAg values >ULOQ (1400 IU/mL) were imputed with 1540 IU/mL; −, negative; +, positive; HBcrAg, hepatitis B core-related antigen; HBeAg, hepatitis B e-antigen; HBsAg, hepatitis B surface antigen; HBV, hepatitis B virus; IU, international unit; LLOQ, lower limit of quantification; NA, nucleos(t)ide analogue; NCT, not currently treated at study start; TND, target not detected; ULOQ, upper limit of quantification; VS, virologically suppressed.
In NCT patients who are HBeAg positive, JNJ-56136379 75 mg and 250 mg + NA resulted in mean (SE) HBeAg declines from baseline of 0.49 (0.09) and 0.70 (0.20) log$_{10}$ IU/mL, respectively, versus 0.81 (0.28) for placebo + NA at Week 24 (table 2; figure 3B; online supplemental figure S4). For VS patients who are HBeAg positive, mean reductions in HBeAg levels were smaller. In patients who are HBeAg positive, the proportion with a >0.3 log$_{10}$ IU/mL reduction in HBeAg from baseline at Week 24 for NCT patients was higher in the combined JNJ-56136379 + NA treatments arms than for placebo + NA (19/23 (83%) vs 5/8 (63%), respectively) and also in VS patients (6/19 (32%) vs 1/5 (20%), respectively) suggesting a trend of more consistent declines across patients (table 2).

Observed declines in HBeAg at Week 24 mostly correlated with declines in HBsAg and early on-treatment isolated alanine aminotransferase (ALT) elevations (ALT flares) (figure 3; online supplemental figure S5). Maximal individual HBsAg and HBeAg reductions were 1.28 and 1.8 log$_{10}$ IU/mL at Week 24, respectively, in the JNJ-56136379 + NA groups and occurred in the patients with the most pronounced HBV DNA declines at Week 24.
Hepatitis B core-related antigen (HBcrAg) changes from baseline were observed between the treatment arms (table 2; online supplemental figure S6).

**Efficacy post-Week 24 of treatment**

Treatment extension with JNJ-56136379 beyond Week 24 had limited additional effect on viral markers (online supplementary table S4). HBsAg and HBeAg declines predominantly occurred early during treatment, but some NCT patients who are HBeAg positive in the JNJ-56136379 250 mg+NA arm...
Figure 5  Mean change from baseline in HBV DNA and HBV RNA over 24 weeks of treatment (pooled placebo/JNJ-56136379+NA treatment arms). (A) Mean change in HBV DNA in NCT patients by HBeAg status. HBV DNA assessed using Roche COBAS HBV DNA assay; HBV DNA values below the LLOQ (20 IU/mL) target detected or target not detected were imputed respectively with 15 IU/mL or 5 IU/mL. Mean baseline HBV DNA levels by treatment group and by HBeAg status are shown in online supplemental table S2. (B) Mean change in HBV RNA in patients who are HBeAg positive and HBeAg negative by prior treatment. HBV RNA assessed using a quantitative reverse transcription PCR assay. Not detected HBV RNA values were imputed by a value of 5 copies/mL. HBeAg, hepatitis B e-antigen; HBV, hepatitis B virus; IU, international unit; LLOQ, lower limit of quantification; NA, nucleos(t)ide analogue; NCT, not currently treated at study start; PCR, polymerase chain reaction; TND, target not detected; VS, virologically suppressed.
had continued HBsAg declines after Week 24 (figure 4; online supplemental figure S2).

None of the patients achieved HBsAg seroclearance and/or seroconversion through Week 48. Two patients achieved HBeAg seroclearance and seroconversion at Week 48 (one VS patient receiving placebo+NA and one NCT patient in the JNJ-56136379 250 mg monotherapy arm). The patient in the JNJ-56136379 250 mg monotherapy arm also had an HBsAg reduction of 0.6 log10 IU/mL at Week 24 with maximal reduction of 1.0 log10 at the last study visit. The patient in the placebo+NA arm with HBeAg seroclearance did not experience any HBsAg reduction.

HBV RNA levels generally increased in patients who stopped JNJ-56136379 treatment at Week 24 or 48 (online supplemental figure S3B). However, kinetics of HBV RNA differed between populations. Some NCT patients who are HBeAg negative reached a new plateau (=1 log10 lower than baseline) in HBV RNA levels, which was maintained during NA treatment in the follow-up period.

ALT normalisation from baseline over time

Most NCT patients with baseline ALT>upper limit of normal (ULN) in the pooled JNJ-56136379 arms and placebo+NA arm achieved ALT normalisation at the end of treatment (Week 24 and/or 48). Of the nine VS patients with baseline ALT>ULN in the pooled JNJ-56136379 arms, four (44%) achieved ALT normalisation versus one (33%) in the pooled placebo+NA arm.

Viral breakthrough

Viral breakthrough (confirmed >1 log10 IU/mL increase in HBV DNA from nadir) occurred in 5/28 patients receiving JNJ-56136379 75 mg monotherapy between Weeks 16 and 20. All five patients had emerging resistant variant T33N (JNJ-56136379-fold change (FC) in EC_{10}~85 as site-directed mutant in in vitro replication assay). NA treatment was initiated in two patients resulting in HBV DNA suppression. The other three patients discontinued the study without follow-up data available. One patient in the JNJ-56136379 75 mg monotherapy arm experienced a viral breakthrough at Week 32, with emerging F23Y mutation (FC=5.2) and was switched to NA treatment with subsequent DNA decline.

One patient receiving JNJ-56136379 250 mg monotherapy with no HBV DNA response up to Week 4 experienced a viral breakthrough at Week 8 with no emerging variants. Following viral breakthrough, NA treatment was added for all patients receiving JNJ-56136379. There were no cases of viral breakthrough in the JNJ-56136379 75 mg+NA or 250 mg+NA treatment arms until Week 24. One patient receiving JNJ-56136379 75 mg+NA experienced a viral breakthrough at Week 48 due to lack of adherence.

Seven patients (six JNJ-56136379±NA and one placebo) experienced a viral breakthrough during the NA-only treatment in the follow-up phase. Of seven patients with a viral breakthrough, three continued ongoing NA treatment and showed subsequent declines in HBV DNA. For the remaining four patients, the time of a viral breakthrough was the last available follow-up time point. No patients had emerging NA-resistance variants. One patient, who received ETV during NA-only follow-up, had documented prior failure to lamivudine and telbivudine and had the M204I lamivudine/telbivudine resistance mutation. Patients with a viral breakthrough and follow-up data available continued NA treatment with subsequent HBV declines.

Safety

Most treatment-emergent AEs were Grade 1 or 2. Three patients discontinued treatment due to AEs (table 3), one in the pooled placebo+NA arm with abdominal discomfort/gastrointestinal upset, one in the JNJ-56136379 75 mg+NA arm with streptococcal toxic shock syndrome, acute cardiac failure, myocarditis and muscle necrosis and one with weight loss in the JNJ-56136379 250 mg open-label arm. Seventeen patients had ≥Grade 3 AEs (13% in the pooled JNJ-56136379 treatment versus four (9%) in the pooled placebo+NA arms), most commonly increased ALT and aspartate aminotransferase (AST) (six (3%) and five (3%) patients, respectively) in the pooled JNJ-56136379 treatment arms versus no patients in the pooled placebo+NA arm (table 3). None of the eight reported SAEs were considered related to study treatment (table 3).

Creatine phosphokinase elevation was reported as an AE in seven (4%) patients in the pooled JNJ-56136379 treatment arms and two (3%) patients in the placebo+NA arm. One patient who received JNJ-56136379 75 mg+NA and one patient who received placebo+NA reported a Grade 3 and Grade 2 AE of creatine phosphokinase elevation, respectively. All other AEs of creatine phosphokinase elevation were Grade 1; cases resolved without discontinuation of treatment.

ALT elevations occurred in seven patients in the JNJ-56136379 treatment arms, including one patient in the JNJ-56136379 75 mg+NA arm (Grade 4), two in the JNJ-56136379 250 mg monotherapy arm (Grade 3) and four in the JNJ-56136379 250 mg+NA arm (Grade 4: n=2; Grade 2: n=1; Grade 1: n=1). These ALT elevations normalised on continued treatment and did not lead to liver failure. No Grade 3 or 4 ALT and AST elevations were observed in the pooled placebo+NA arm (online supplemental table S5). In the pooled JNJ-56136379 treatment arms, transient ALT elevations of Grade 3 were observed in 10 (5%) patients and of Grade 4 in five (3%) patients. Transient AST elevations of Grade 3 were observed in three (2%) patients and of Grade 4 in two (1%) patients.

There were two patients with peak ALT >1000 UL (1340 and 1086 UL) from the JNJ-56136379 250 mg+NA and JNJ-56136379 75 mg open-label arms, respectively. An additional 10 patients had peak ALT values between 300–1000 UL (JNJ-56136379 75 mg open-label: n=2; JNJ-56136379 250 mg open-label: n=3; JNJ-56136379 75 mg+NA: n=1; JNJ-56136379 250 mg+NA: n=4). Most ALT flares occurred around Weeks 1 and 2, and time until resolution was typically between 2 and 3 weeks while treatment was continued.

Patients with estimated glomerular filtration rate based on serum creatinine (eGFR_u) >60 mL/min/1.73 m² (lower limit of Grade 2) or higher were included in the study and sometimes declined to <60 mL/min/1.73 m², classified as Grade 3 decreases. An on-treatment decline in eGFR_u was observed early after JNJ-56136379+NA treatment initiation with rapid increase during the follow-up phase. Figure 6 shows fluctuations in eGFR_u over time, by type of NA. The reductions appeared to be more pronounced in patients receiving JNJ-56136379 250 mg±NA versus JNJ-56136379 75 mg±NA. A higher proportion of patients in the JNJ-56136379 250 mg±NA group had treatment emergent Grade 3 eGFR_u during study treatment (online supplemental...
The decline in eGFR\textsubscript{cr} was observed at the first assessment after starting JNJ-56136379. The decline in eGFR\textsubscript{cr} stabilised at subsequent assessments, and there was a rapid increase in eGFR\textsubscript{cr}. A conclusion cannot be drawn regarding the type of NA and impact on eGFR\textsubscript{cr} given the small sample size of patients receiving ETV.
Pharmacokinetics

After repeated qd dosing with JNJ-56136379 75 mg and 250 mg up to 24 weeks, observed plasma concentrations were dose proportional and within the range predicted from the population PK model (online supplemental figure S7A). The PK data were well described using a two compartmental model (central and peripheral distribution volume). Body weight was retained as a covariate to explain the volumes, including sex-specific reference weights, but age was not retained as a covariate in the model. An evaluation of the PK data from JNJ-56136379 indicated that the initial population PK model (using the same approach for body weight and the same parameters) generally well described the JNJ-56136379 concentrations observed in this study for both the 75 mg and 250 mg doses in the initial and extension treatment phases. No additional effects of race (Asian vs non-Asian) were observed.

Comparing plasma concentrations and PK parameters between the JNJ-56136379 monotherapy arms and combination arms with an NA, a PK drug-drug interaction between TDF and JNJ-56136379 was apparent. At Day 84, tenofovir disoproxil fumarate (TFV) exposure was higher for both arms containing JNJ-56136379 versus placebo (online supplemental figure S7B).

In comparison with arms receiving placebo at Week 12, for all arms receiving JNJ-56136379 75 mg and 250 mg, respectively, the geometric mean ratios of TFV maximum plasma concentrations (Cmax) were 142.81% and 165.02%, TFV included pre-dose plasma concentrations (Ctough) were 149.21% and 171.74% and TFV area under the plasma concentration-time curve from time 0 to τ hours post-dose (AUCτ) were 189.55% and 195.06%. Drug-drug interactions between ETV and JNJ-56136379 were not apparent (online supplemental figure S7C).

DISCUSSION

While the primary endpoint of >1 log10 IU/mL mean decline in HBsAg of the active versus control was not achieved, this study demonstrated that JNJ-56136379 75 mg and 250 mg qd administered in combination with TDF or ETV led to substantial reductions in HBV DNA and HBV RNA, up to Week 48, and had favourable safety and tolerability profiles. For NCT patients who are HBeAg positive, a trend of a greater, yet small HBsAg reduction in plasma concentrations (Cmax) was 142.81% and 165.02%, TFV included pre-dose plasma concentrations (Ctough) were 149.21% and 171.74% and TFV area under the plasma concentration-time curve from time 0 to τ hours post-dose (AUCτ) were 189.55% and 195.06%. Drug-drug interactions between ETV and JNJ-56136379 were not apparent (online supplemental figure S7C).

As reported previously, JNJ-56136379 showed potent reductions of serum HBV RNA. In NCT patients, a greater mean HBV RNA decline was observed in the JNJ-56136379 75 mg+NA and 250 mg+NA groups versus placebo (2.96 and 3.15 versus 1.33 log10 copies/mL) at Week 24. In VS patients, mean HBV RNA declines appeared lower than in NCT patients due to low baseline HBV RNA levels and the high proportion of patients with already undetectable HBV RNA at baseline. Serum HBV RNA has recently been suggested as a measure for cccDNA transcriptional activity. However, its use as a marker for cccDNA is limited in the setting of CAM treatment given their direct effect (primary MoA), preventing pgRNA encapsidation and thereby, release of HBV RNA-containing particles; thus, in the setting of CAM administration, serum HBV RNA is primarily

the reflection of target engagement. A rebound in HBV RNA after stopping JNJ-56136379 under continued NA treatment was observed, suggesting mainly a direct effect on HBV RNA release.

HBcrAg and HBeAg are derived from cccDNA and considered surrogate markers for cccDNA transcriptional activity. If a possible effect of JNJ-56136379+NA treatment on cccDNA occurred, it may be reflected in HBeAg and HBcrAg levels.

In the present study, compared with NA only, JNJ-56136379+NA had limited additional effects on HBeAg (such as more patients with >0.3 log decline), which may be due to effects on cccDNA or direct effects of the CAM-N on HBeAg secretion. However, no relevant effect on HBcrAg levels was noted.

As a viral breakthrough associated with resistant variants was observed in the JNJ-56136379 monotherapy arm, JNJ-56136379 will not be developed further. These data imply that the risk of viral resistance and breakthrough should be considered for the drug class of CAMs in general. The JNJ-56136379 emerging variant was detectable as the dominant variant until the last visit with sequence data available. The analysis of persistence of CAM-related variants after stopping JNJ-56136379 is hampered by the fact that either NA rescue treatment was started in most patients resulting in subsequent HBV DNA declines or the patients were lost to follow-up. No cases of viral breakthrough were observed in the JNJ-56136379+NA combination arms through Week 24, although one patient receiving JNJ-56136379 75 mg+NA experienced viral breakthrough at Week 48, likely due to adherence issues.

JNJ-56136379 was well tolerated, where most treatment-emergent AEs were Grade 1 or 2; no study treatment-related SAEs were observed. Transient Grade 3 and 4 ALT/AST elevations without signs of impaired liver function were observed in the JNJ-56136379+NA arm. ALT/AST elevations normalised on continued treatment. ALT flares were primarily observed in NCT patients who are HBeAg positive and were often associated with decreases in viral parameters. Isolated and transient treatment-emergent ALT and AST increases have been observed during treatment with NAs, mainly in patients who are HBeAg positive.

JNJ-56136379±NA treatment caused a transient decrease in eGFR, with an increase during the follow-up phase. The increase in plasma creatinine that occurred shortly after treatment initiation with JNJ-56136379 suggests a renal transporter effect of JNJ-56136379 at the proximal tubular level, leading to decreased tubular excretion of creatinine and lower eGFR

values. Plasma levels of JNJ-56136379 were dose-proportional as reported previously. There was no evidence of a drug-drug interaction between JNJ-56136379 and ETV. TFV exposure was twofold higher in combination with JNJ-56136379 versus placebo. Although these changes were not considered clinically relevant, the clinical impact of longer treatment exposure to a combination regimen of JNJ-56136379 and TDF is unknown. Therefore, further studies are needed to evaluate if the increase in TFV exposure is unique to co-administration with JNJ-56136379 or if similar increases are observed for the CAM class.

Strengths of the study were the comprehensive design that included a blinded comparison of active versus placebo, NCT and VS patients, HBeAg-positive and -negative patients, relevant numbers of Asian patients and patients from China, mono-therapy and combination treatment, two doses of study drug and a response guided longer treatment period to better characterise antiviral activity. Limitations included the low power to fully assess the primary and secondary endpoints in patient subgroups
and the initial treatment extension criteria with a stringent threshold for HBV DNA at 24 weeks, which may have excluded patients who could have responded with further treatment.

In conclusion, treatment with JNJ-56136379 in combination with an NA in patients with CHB showed pronounced reductions in HBV DNA and HBV RNA. Limited effects of JNJ-56136379 on HBSAg and HBeAg levels were observed. Both doses of JNJ-56136379 in combination with an NA were well tolerated over a prolonged treatment period and demonstrated target engagement. While JNJ-56136379 in combination with an NA did not show a clear benefit over NA monotherapy, several other CAMs of different classes and chemotypes, with improved preclinical characteristics are in discovery and in clinical development. The results of these studies will inform on the role of CAMs in future combination regimens for the treatment of HBV.

**Author affiliations**

1Toronto General Hospital, Toronto, Ontario, Canada
2Erasmus Medical Center, Rotterdam, Zuid-Holland, The Netherlands
3Nanfang Hospital, Southern Medical University, Guangzhou, China
4Université de Paris Cité, INSERM UMR1149, Hôpital Beaujon AP-HP, Clichy, France
5The Chinese University of Hong Kong, Hong Kong SAR, China
6Hospices Civils de Lyon and Lyon University & INSERM U1052-Cancer Research Institute of Lyon, Lyon, France
7Department of Gastroenterology and Hepatology, Kumamoto University, Kumamoto, Japan
8Faculty of Health Sciences in Bytom, Medical University of Silesia, Katowice, Poland
9JD Care, Hillsborough, New Jersey, USA
10ZNA Jan Pallijin, CPU, Antwerp, Belgium
11Hospital Universitario Vall d’Hebron and CIBERHED del Instituto Carlos III, Barcelona, Spain
12Division of Gastroenterology and Hepatology, Foundation IRCCS Ca’ Granda Ospedale Maggiore Policlinico, Milan, Italy
13Department of Pathophysiology and Transplantation, CRC ‘A. M. and A. Migliavacca’ Center for Liver Disease, University of Milan, Milan, Italy
14Janssen Pharmaceutica NV, Beerse, Belgium
15Janssen Pharmaceuticals R&D, Titusville, New Jersey, USA

**Acknowledgements**

The authors would like to thank the patients and their families, the study investigators and site staff and all the members of the JNJ-56136379 Compound Development and Operations Teams for conducting this study and other Janssen staff members for their important contributions to the manuscript. Medical writing support for the development of this manuscript was provided by Ian Woodever of Ashfield MedComms, an Ashfield Health company, and Jessica Swanner, PhD, of Luminary Communications Inc., and was funded by Janssen.

**Contributors**

HLAJ, JH, TA, HLYC, FZ, YT, EJ, RGN, SB, MBu and PL contributed to the conduct of the studies as investigators and to the interpretation of the data. OL, TV, JT, WT, MBe, RK, MBI and US contributed to the design and conduct of the study, analysis and interpretation of the data. All authors were involved in manuscript development, review and revision, interpretation of data and have read and approved the final version to be submitted. All authors satisfy the criteria for authorship as established by the ICMEI. All authors had access to the study data and reviewed and approved the final manuscript. MBI served as the guarantor of the study.

**Funding**

This study was sponsored by Janssen (award/grant number: not applicable).

**Competing interests**

HLAJ received grants from AbbVie, Arbutus, Gilead Sciences, Janssen and Roche, and is a consultant for Arbutus, Arena, Enyo Pharma, Gilead Sciences, GlaxoSmithKline, Janssen, Merck, Roche, Vir Biotechnology and Vicriviroc. JH declares no conflicts of interest. TA has received grants and served as a speaker and clinical investigator for AbbVie, Antios Therapeutics, Enyo Pharma, Eiger BioPharmaceuticals, Gilead Sciences, Janssen, Merck and Roche. HLYC is an advisor for AbbVie, Alios, Arbutus, Gilead Sciences, Hepion, Janssen, Merck, Roche, Vir Biotechnology, Vicriviroc and Viroceutics, and served as a speaker for Gilead Sciences, Mylan and Roche. FZ has participated in advisory committees or review panels for Janssen, Gilead Sciences, AbbVie, Arbutus, Transgene, Contravir, Mypharma, Spring Bank, Algos and Assembly, received grant/research support from Roche and Sanofi/Epocet, and speaking and teaching support from Gilead Sciences. YT received lecture fees from Fujirebio, Symsysx and Gilead Sciences, conceived/joint research agreements from Fujirebio, Janssen, Gilead Sciences, GlaxoSmithKline and Stanford Junior University, and has participated in advisory boards for Gilead Sciences and GlaxoSmithKline. EJ has served as a speaker and clinical investigator for AbbVie, Bristol Myers Squibb, Gilead Sciences, Janssen, Merck and Roche. RGN has received grant/clinical research support from Janssen, Assembly, AbbVie, Alkermes, Gilead Sciences, Merck and Viiv, and has participated in speaker panels for Gilead Sciences, Merck, Insemed and Viiv. SB has participated in advisory committees or review panels for AbbVie, Gilead Sciences, Merck, Roche and Spring Bank Board, board membership for AbbVie, received grant/research support from Gilead Sciences and Janssen, and speaking and teaching support from Gilead Sciences, Janssen and Bristol Myers Squibb. PL has participated in advisory boards/speaker bureaus for Bristol Myers Squibb, Roche, Gilead Sciences, GlaxoSmithKline, AbbVie, Merck, Arrowhead, Ahylam, Janssen, Spring Bank Board, Mypharma and Eger. OL, TV, JT, WT, MBe, RK, MBI and US are all employees of Janssen Pharmaceuticals and Johnson & Johnson stockholders.

**Patient and public involvement**

Patients and/or the public were not involved in the design, or conduct, or reporting, or dissemination plans of this research.

**Patient consent for publication**

Not applicable.

**Ethics approval**

This study involves human participants and was approved by the relevant local institutional review boards/ethics committees. Participants gave informed consent to participate in the study before taking part.

**Provenance and peer review**

Not commissioned; externally peer reviewed.

**Data availability statement**

The data sharing policy of Janssen Pharmaceutical Companies of Johnson & Johnson is available at https://www.janssen.com/cclinical-trials/transparency. As noted on this site, requests for access to the study data can be submitted through Yale Open Data Access (YODA) Project site at http://yoda.yale.edu.

**Supplemental material**

This content has been supplied by the author(s). It has not been vetted by BMJ Publishing Group Limited (BMJ) and may not have been peer-reviewed. Any opinions or recommendations discussed are solely those of the author(s) and are not endorsed by BMJ. BMJ disclaims all liability and responsibility arising from any reliance placed on the content. Where the content includes any translated material, BMJ does not warrant the accuracy and reliability of the translations (including but not limited to local regulations, clinical guidelines, terminology, drug names and drug dosages), and is not responsible for any errors and/or omissions arising from translation and adaptation or otherwise.

**Open access**

This is an open access article distributed in accordance with the Creative Commons Attribution Non Commercial (CC BY-NC 4.0) license, which permits others to distribute, remix, adapt, build upon this work non-commercially, and license their derivative works on different terms, provided the original work is properly cited, appropriate credit is given, any changes made indicated, and the use is non-commercial. See: http://creativecommons.org/licenses/by-nc/4.0/.%0A

**ORCID iDs**

Jinlin Hou http://orcid.org/0000-0001-8230-8583
Fabián Zoulím http://orcid.org/0000-0002-2245-0083
Pietro Lampertico http://orcid.org/0000-0002-1026-7476
Michael Biermer http://orcid.org/0000-0001-6993-8206%0A

**REFERENCES**


Hepatology

Gut first published as 10.1136/gutjnl-2022-328041 on 25 January 2023. Downloaded from http://gut.bmj.com/ on September 23, 2023 by guest. Protected by copyright. %0A


12 Zoulim F, Lenz O, Vandenbossche JJ, et al. JNJ-56136379, an HBV capsid assembly modulator, is well-tolerated and has antiviral activity in a phase 1 study of patients with chronic infection. Gastroenterology 2020;159:521–33.

13 Gane E, Schwabe C, Lenz O. JNJ-64530440 (JNJ-0440), a novel class N capsid assembly modulator (CAM-N): safety, tolerability, pharmacokinetics (PK), and antiviral activity of multiple ascending doses in patients (PTS) with chronic hepatitis B (CHB). Hepatology 2019;70.


