

Will we need novel combinations to cure HBV?

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Disclosures

- Gilead Sciences
- Janssen
- Arbutus

Combination treatment: Evidence still being gathered

- Limited clinical data to date
 - Several steps in the replication cycle of HBV are druggable targets
 - Safety of new combinations will be paramount
 - (Given the safety of currently approved nucleoside analogues).
-

Strategies for HBV eradication



TARGETING HBSAG



TARGETING
INTEGRATED VIRAL
DNA



HBX PROTEIN



STRATEGIES TO
INTERFERE WITH
VIRAL ENTRY



CCCDNA
REGULATION



RNA INTERFERENCE
AGENTS



INNATE IMMUNE
AGONISTS



IMMUNE
THERAPIES

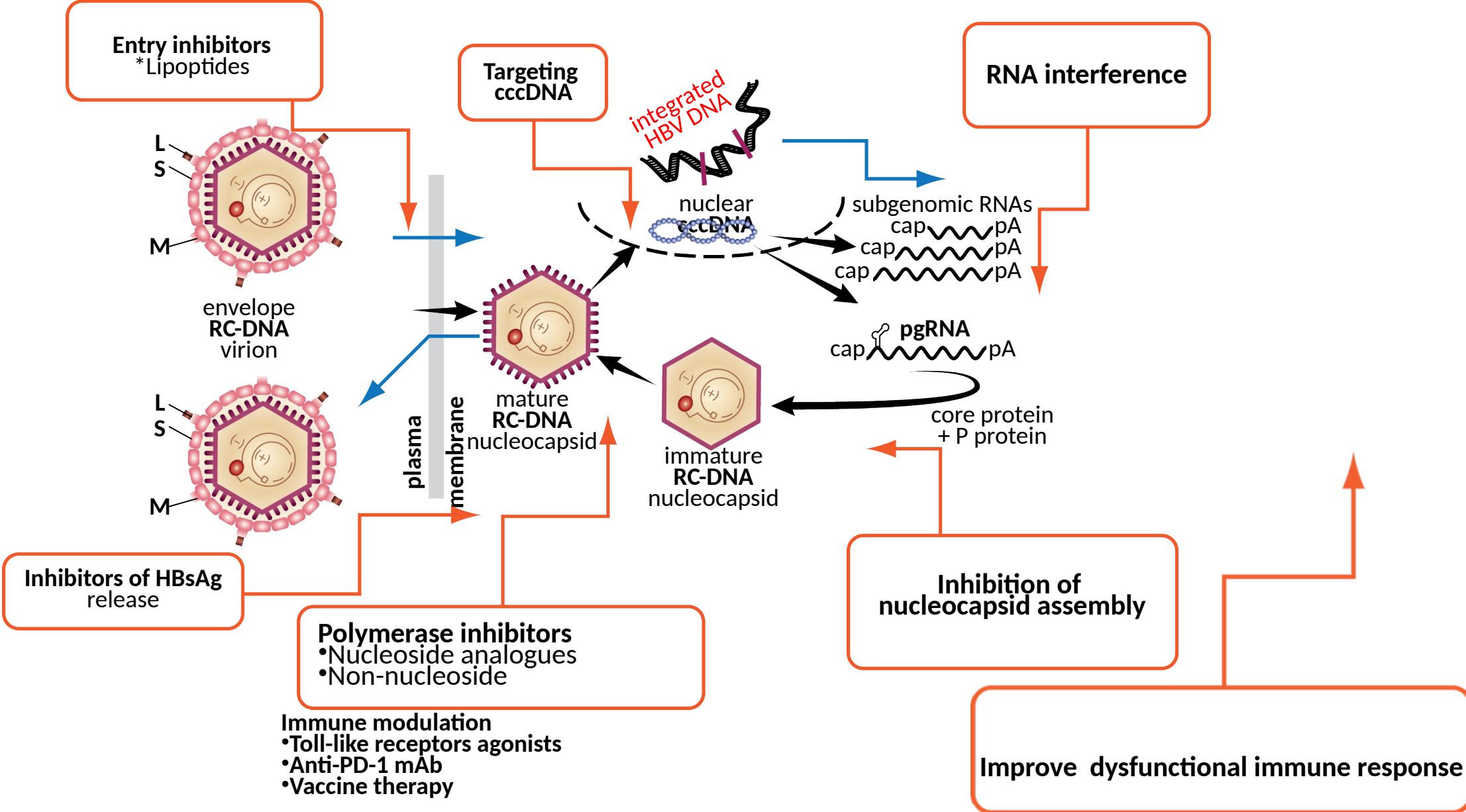


CAPSID ASSEMBLY
MODULATORS

Virology of HBV poses difficulties for cure

- HBV DNA genomic fragments integrate into the genome of hepatocytes
- The stable episomal cccDNA acts as the template for transcription of HBV mRNAs
- HBsAg can be transcribed from cccDNA and integrated viral genomes
 - HBeAg negative patients the probable predominant source of HBsAg is from RNAs transcribed from integrant HBV DNA
 - This source of HBsAg is relatively inaccessible without cell loss or DNA editing
- In addition to infectious virions, HBV replication results in the excess production and release of sub-viral empty envelope particles
- HBsAg antigen load may cause profound antigen-specific immune dysfunction and exhaustion.
- Selection of resistant associated substitutions possible with eg CAMs

HBV antiviral targets: exploitable targets





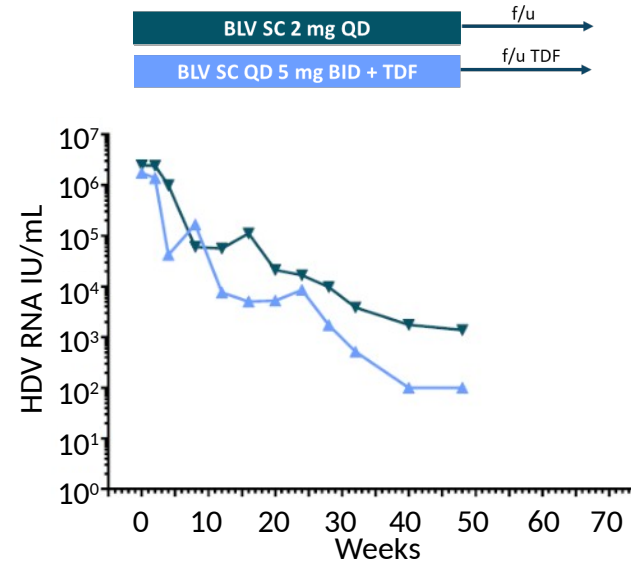
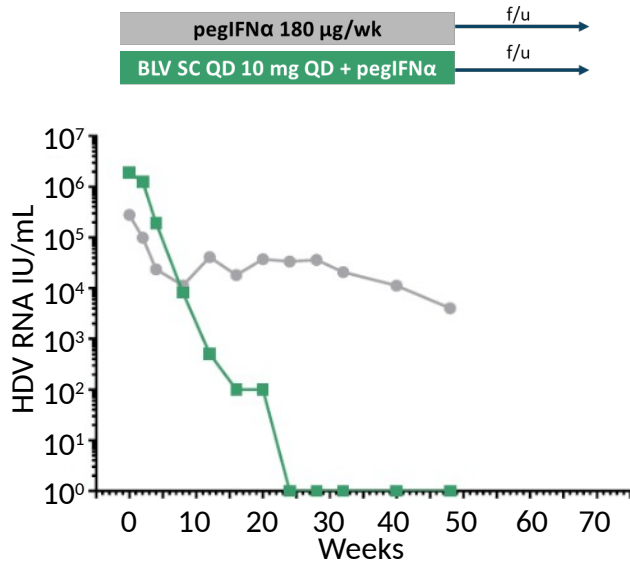
Hepatitis D infection

10 mg bulevirtide (Myrcludex B) in combination with pegIFN-α2a or tenofovir

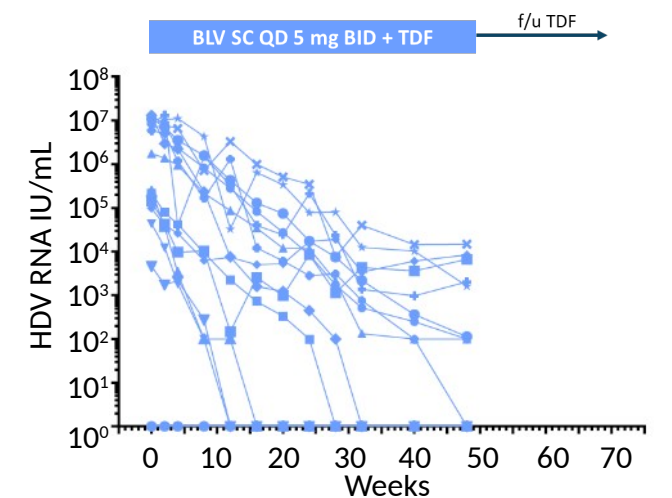
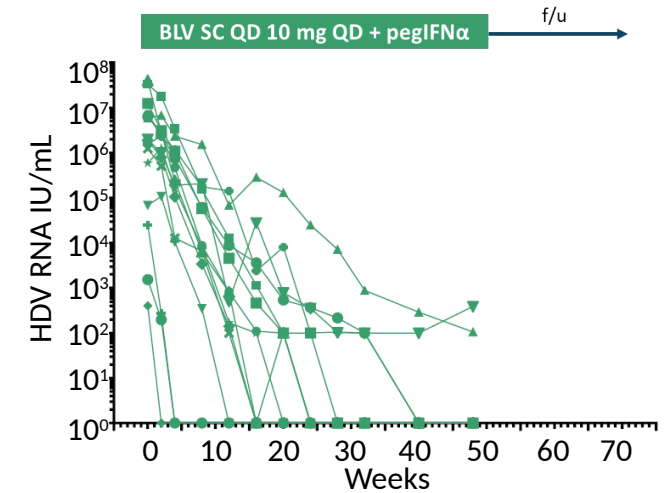
Chronic HBV/HDV coinfection: Week 24 interim results of the MYR203 extension study



Virologic response (median HDV RNA)



Individual RNA kinetics



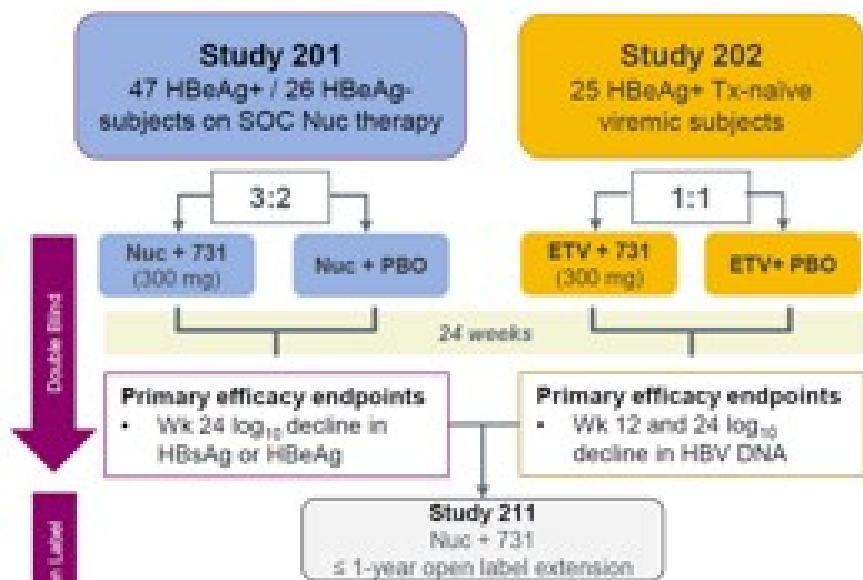
Virologic response: Week 48	Median HDV RNA reduction (log)	Undetectable HDV RNA
pegIFNα	-1.29	13.3%
2 mg BLV + pegIFNα	-5.21	80.0%
5 mg BLV + pegIFNα	-6.13	86.7%
10 mg BLV + pegIFNα	-6.09	86.7%

Virologic response: Week 48	Median HDV RNA reduction (log)	Undetectable HDV RNA
2 mg BLV	-2.84	13.3%
5 mg BLV BID + TDF	-4.58	40.0%



HBV capsid assembly modulators

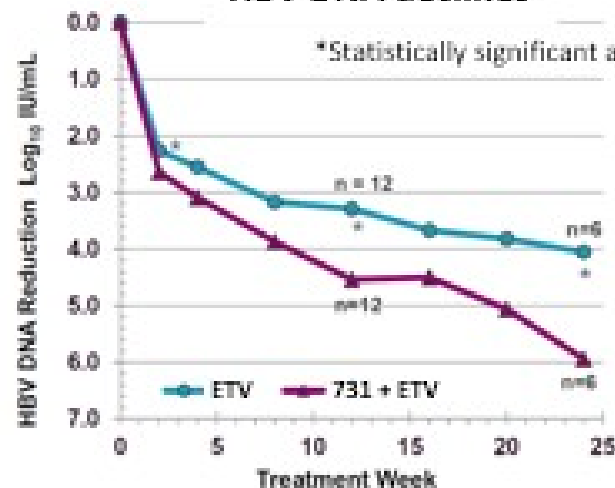
Interim safety and efficacy results of ABI-H0731 Phase 2a program exploring combination of ABI-H0731 with Nuc therapy in treatment-naïve and treatment-suppressed CHB patients



Blinded summary of TEAEs (Studies 201 and 202)

- No SAEs or treatment related d/cs or interruptions
- AEs mostly mild, infrequent, considered unrelated to study drug
- No flares on treatment; no clinical AE ≥ Grade 2
- 3 pts with rash considered “possibly related” (2x Gr 1, 1x Gr 2)
- 1 pt in each study had Gr 2 AE considered possibly related to drug
 - Macular/maculopapular rash—resolved on antihistamine (201)
 - ALT increase—resolved with continued treatment (202)

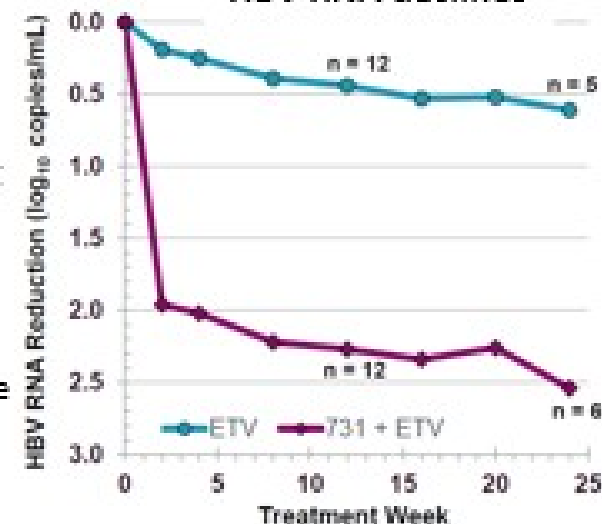
HBV DNA declines



Week	ETV	ETV + 731	p-value
12	3.29	4.54	<0.011
24	3.99	5.94	<0.005

HBV DNA assessed by Roche CobasqPCR; LOQ = 20 IU

HBV RNA declines

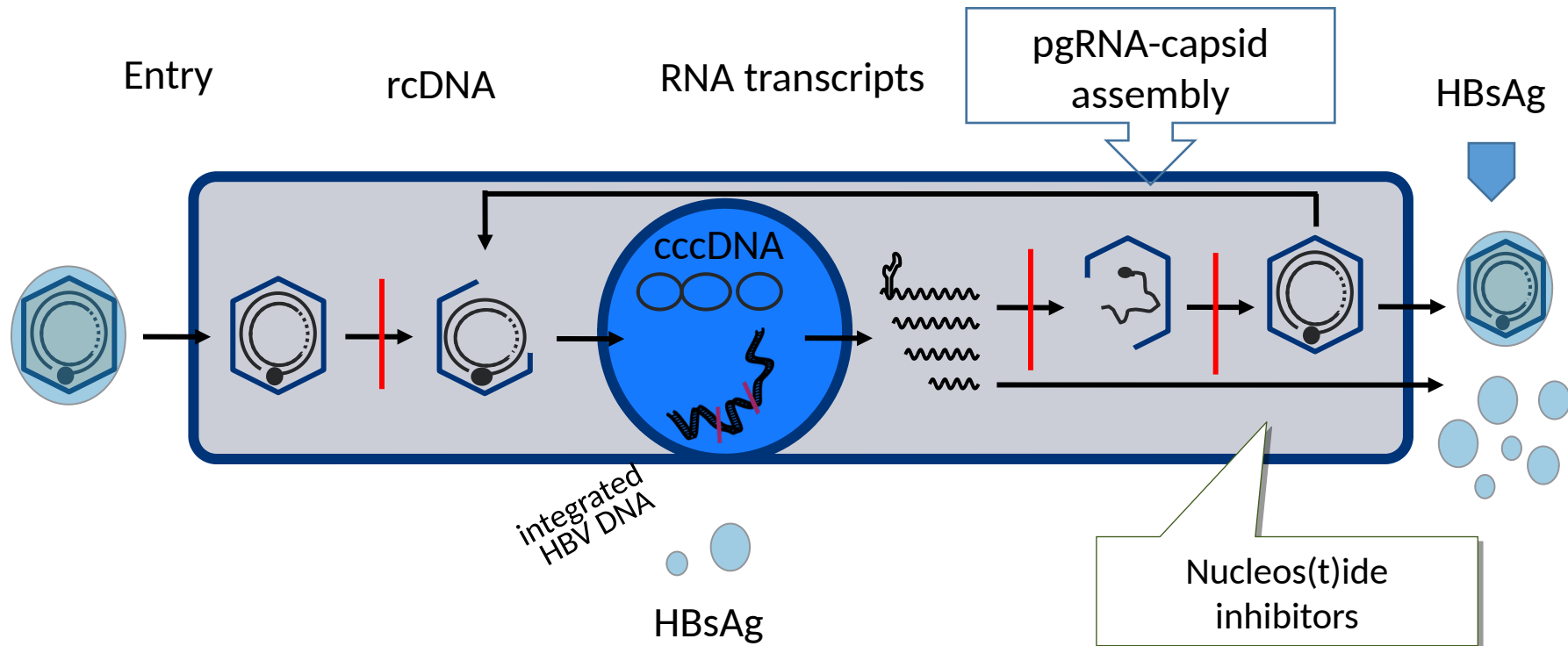


Week	ETV	ETV + 731	p-value
12	0.44	2.27	<0.005
24	0.61	2.54	<0.005

HBV RNA assessed by RT qPCR; LOQ = 200 copies/mL

Capsid Assembly modifier + Nucleoside analogues: effect on HBV replication: effect on cccDNA and HBsAg?

Continued transcription from cccDNA and integrated viral genomes: relatively minor decrease serum HBsAg despite undetectable serum HBV DNA



rcDNA – relaxed circular DNA; cccDNA –covalently closed DNA; pgRNA pre-genomic RNA



RNA therapeutics: RNAi

RNAi-based triple combination therapy in CHB: JNJ-3989 (RNAi), JNJ-6379, (CPAM), and nucleoside analogue



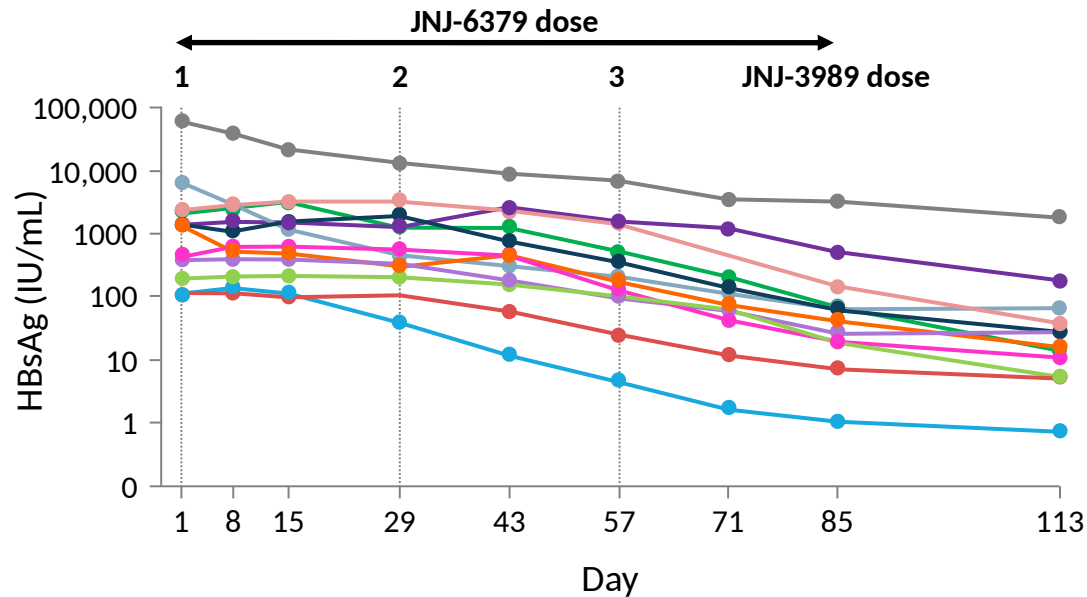
Background

JNJ-3989 Q4W 3 monthly 100–400 mg with Nuc >1.0 log drop HBsAg and reduced all measurable viral products (EASL 2019)

JNJ-6379 blocks HBV replication and *de novo* cccDNA pathway achieving significant decline in serum HBV DNA and serum HBV RNA (AASLD 2018)

Aim

To help design longer term studies, a cohort added to achieve experimental triple combination of JNJ-3989, JNJ-6379, and NA

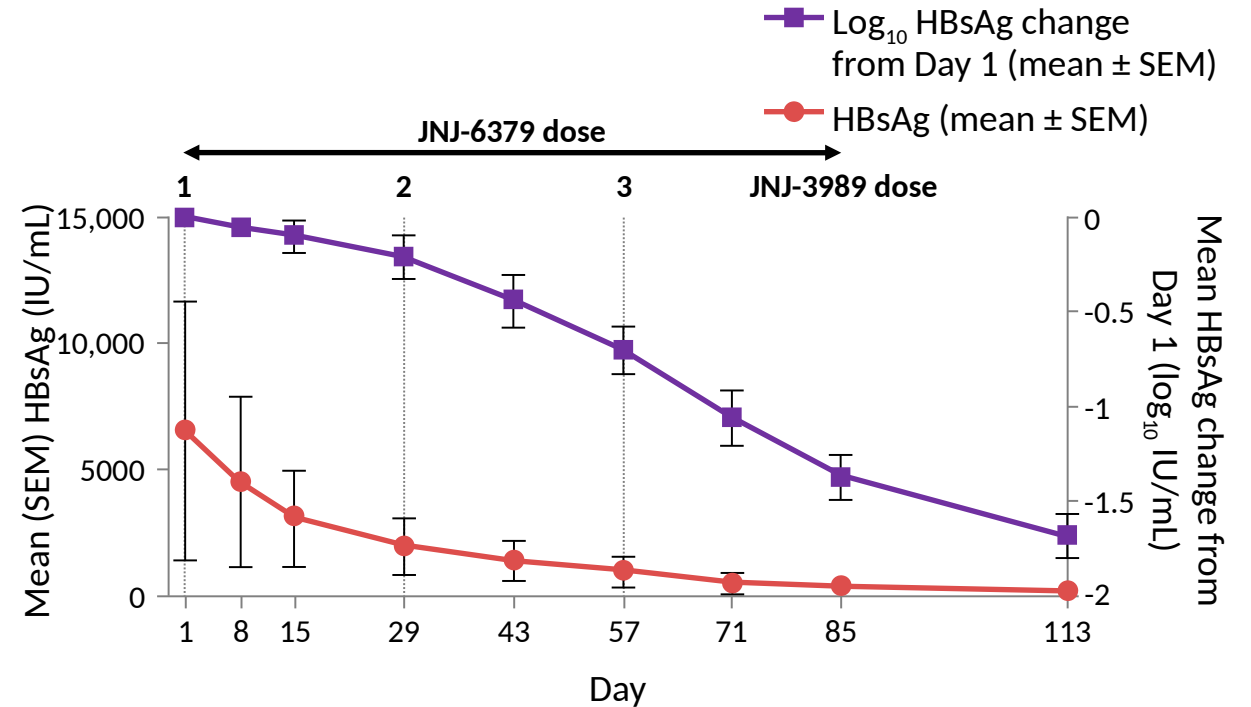


Patients

- 12 patients; 3 doses JNJ-3989 200 mg Day 1, 29, and 57 SC + oral JNJ-6379 250 mg OD for 12 weeks

Results

- Efficacy data to Day 113 (2 months post-3898 and 1 month post-6378)





IHEP HBV

Antisense oligonucleotides (ASOs)

12 weeks' treatment of multiple doses of GSK3389404 (ASO) in CHB subjects on stable nucleoside therapy

Phase 2a, double-blind, placebo-controlled study



GSK3389404: 2nd-generation liver-targeted antisense oligonucleotide (ASO) that targets all HBV-derived RNA transcripts

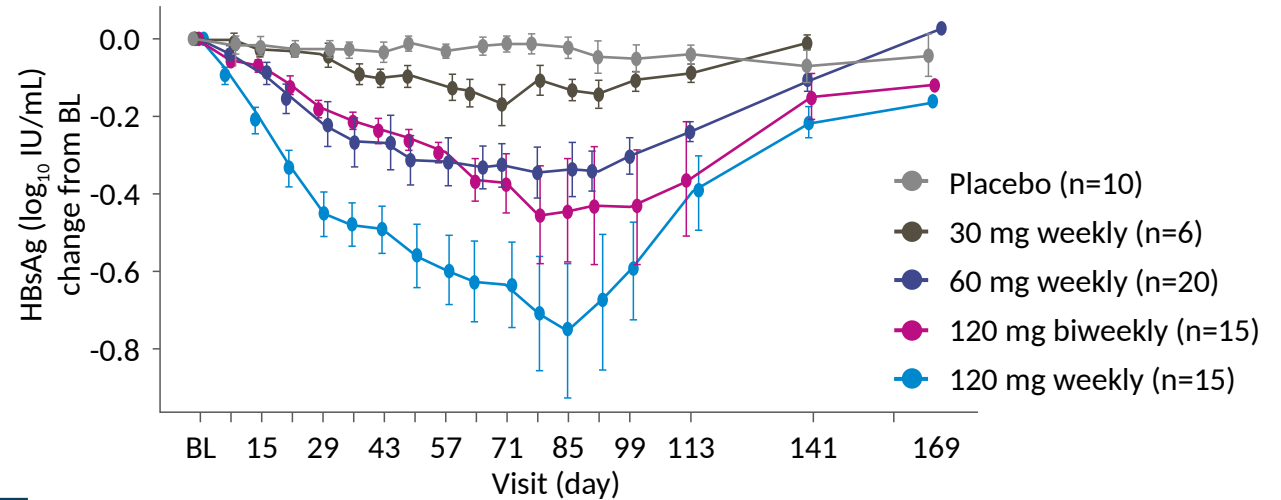
Phase 2a, multicenter, randomized, double-blind, placebo-controlled study in Asia-Pacific region

Multiple-dose, 12 weeks SQ injection; study arms:

- 60 mg weekly
- 120 mg bi-weekly
- 120 mg weekly
- PBO

HBeAg+/- Nuc-treated non-cirrhotic CHB patients with HBsAg >50 IU/mL and ALT ≤2 x ULN

Mean change from BL in HBsAg (log₁₀ IU/mL) over time



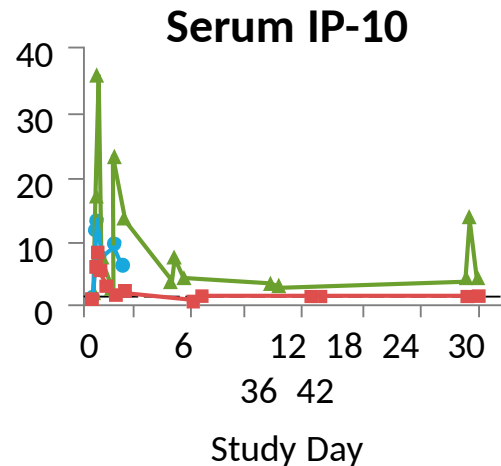
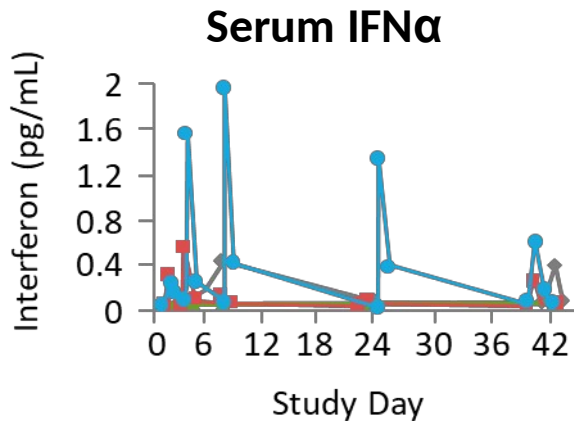
- Dose-dependent declines in mean HBsAg
 - Similar in HBeAg+ and HBeAg- subjects
 - 3 subjects achieved ≥1.5 log₁₀ decrease in HBsAg with no ALT flare
 - 1.54 log₁₀ in 60 mg weekly arm on Day 85
 - 2.37 log₁₀ in 120 mg biweekly arm on Day 92
 - 2.72 log₁₀ in 120 mg weekly arm on Day 85
- Mean HBsAg declines of 0.02 log IU/mL in PBO, 0.13 log IU/mL in 30 mg weekly, 0.34 log IU/mL in 60 mg weekly, 0.44 log IU/mL in 120 mg biweekly, and 0.75 log IU/mL in 120 mg weekly treatment arms by Day 85

	PBO	30 mg weekly	60 mg weekly	120 mg biweekly	120 mg weekly
N	10	6	20	15	15
HBeAg+	30%	17%	40%	13%	27%
Mean HBsAg log ₁₀ , IU/mL	3.15	2.73	2.96	2.82	3.00
ALT, U/L	22.9	24.5	18.6	21.1	19.1
Platelets, (10 ⁹ /L)	214	194	243	200	229

TLR-7 agonist RO7020531

- Liver targeting specific TLR-7
 - 150 mg QOD dosing for 6 weeks in NA-suppressed CHB patients

PD activity in patients with flu-like symptoms



Relationship between exposure and PD activity

	Fraction responding	Geometric mean fold change (range)
Neopterin	6/8	3.13 (1.86–6.08)
IP-10	7/8	3.55 (1.37–36.43)
ISG15	8/8	11.21 (2.31–270.26)
OAS-1	8/8	4.85 (1.71–41.45)
MX1	8/8	6.78 (2.16–87.43)
TLR7	7/8	3.46 (2.04–6.84)

- AASLD 2019 additional 150 and 170 mg cohorts (Yuen #692)
 - full virologic results included a HBsAg decline in qHBsAg
 - Next year planned Phase II platform studies with other agents

Gane E, et al. AASLD 2018, San Francisco, USA. #LB-33 Yuen M-F, et al. AASLD 2019, Boston, USA. #692



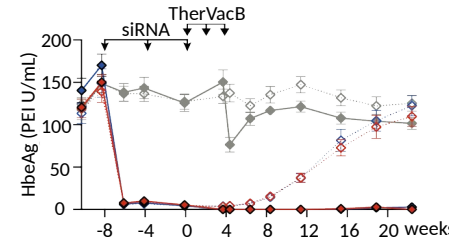
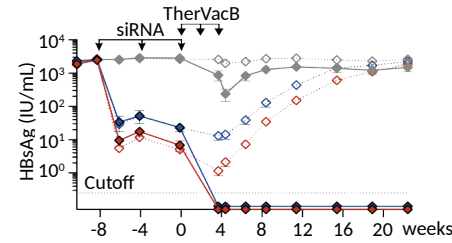
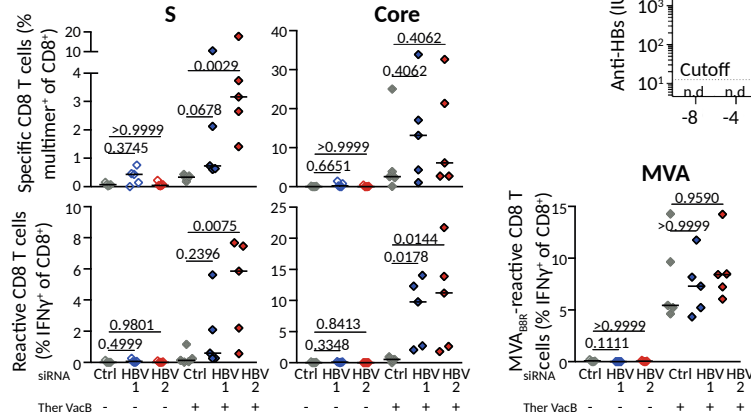
Experimental mouse models

Long-term functional cure in preclinical mouse model

Heterologous therapeutic hepatitis B vaccine (**TherVacB**) consisting of a protein prime / Modified Vaccinia Ankara Virus (MVA) boost vaccination

N-acetylgalactosamine-coupled siRNA enables hepatocyte-specific delivery of siRNA targeting the common 3' end of HBV transcripts to suppress all viral transcripts and proteins

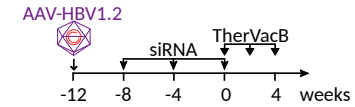
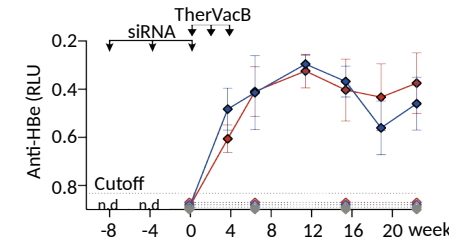
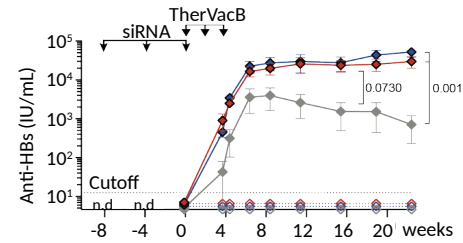
T cell responses (5 months after MVA-boost)



Legend for HBsAg and HbeAg graphs:

- siRNA Ctrl
- siRNA HBV1
- siRNA HBV2

TherVacB: \Rightarrow



AAV-HBV1.2: 2×10^{10} geq i.v.
 GalNAc-siRNAs: 3 mg/kg bw s.c.

n.d. not determined

➤ Functional cure in all 12 mice treated with HBV siRNA \Rightarrow TherVacB
 ➤ 3/12 animals showed mild and transient HBeAg relapse 4 months after stop of treatment

- Combinational siRNA/vaccination therapy achieved functional cure through induction of virus-specific CD8⁺ T cell response
- Proof of concept on suppression of all viral antigens but not HBV DNA alone is needed for immune induction by therapeutic vaccine

Conclusions: Will we need novel combinations to cure HBV?

Synergistic mechanisms need to incorporate a decrease in HBV transcription, loss of cccDNA or altered epigenetic regulation of cccDNA and immune modulation or immunologically stimulated hepatocyte cell turnover.

Nucleoside analogue suppressed patients being included in many current trials. Trials are progressing to combination therapy as additive or synergistic effects are sought.

Trials will provide insights into the biology of HBV and perturbations of the immune response, required to effect HBsAg loss at different stages of the disease.