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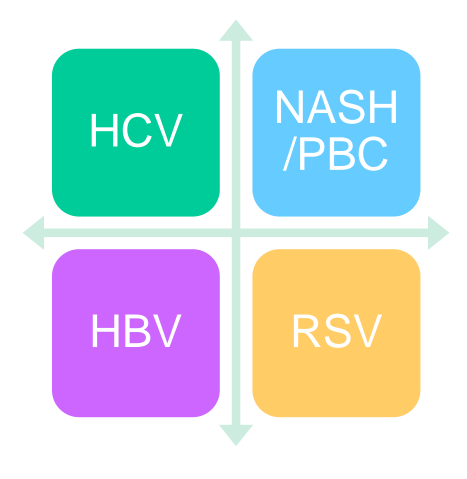
Investment Highlights

- Virology & liver disease-focused biotech company
- Two partnered products marketed in AbbVie's HCV regimens:
 - Glecaprevir HCV protease inhibitor in MAVYRET™/MAVIRET™
 - Paritaprevir HCV protease inhibitor in VIEKIRA* regimens
 - Fiscal 2Q19 royalties on HCV regimens: \$39.6 million
- Three clinical-stage programs in areas of high unmet medical need:
 - RSV: Phase 2a human challenge study complete
 - NASH: Phase 2 "ARGON-1" study ongoing
 - PBC: Phase 2 "INTREPID" study ongoing
- Ongoing R&D programs in NASH/PBC, HBV and RSV
- Strong balance sheet to fund clinical programs and other R&D efforts
 - Approx. \$386.7M in cash at 3/31/19



Our Therapeutic Focus

- Leverage our core strength in HCV to become a leader in Viral and Liver diseases
- Multiple new therapeutic areas with goal of building multiple approaches in each





Broad Virology and Liver Disease Pipeline

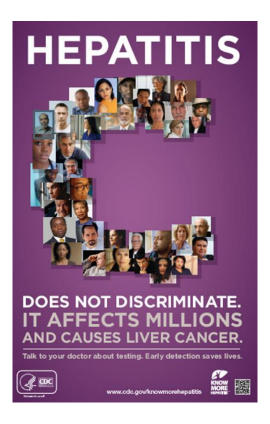
Product Candidate		Discovery	Preclin	Phase 1	Phase	e 2	Phase 3	Market
HCV	Protease Inhibitor	glecaprevir –	glecaprevir – containing pan-genotypic 2-DAA combo				MAVYRET = gleoaprevir/pibrentasvir	
HCV	Protease Inhibitor	paritaprevir -	paritaprevir – containing regimens					
RSV	N-protein Inhibitor	EDP-938	Ph2 C	hallenge Study	/			
NASH	FXR Agonist	EDP-305	Ph2 "A	ARGON-1"				
PBC	FXR Agonist	EDP-305	Ph2 "II	NTREPID"				
HBV	Core Inhibitor	EDP-514						
NASH	FXR Agonist Follow-on							
NASH	Undisclosed							



Glecaprevir – A Pan-genotypic HCV Protease Inhibitor

- Glecaprevir: the protease inhibitor in AbbVie's MAVYRET™*
 - RBV-free, once-daily, fixed-dose combination (2-DAA)
 - MAVYRET treats the majority of patients today in only 8-weeks
- Also treats patients with specific challenges:
 - compensated cirrhosis
 - severe chronic kidney disease
 - PI or NS5A treatment failures
- Marketed by AbbVie (U.S., EU, Japan & other countries globally)
- Market for HCV therapies: ~ \$8 billion in 2018

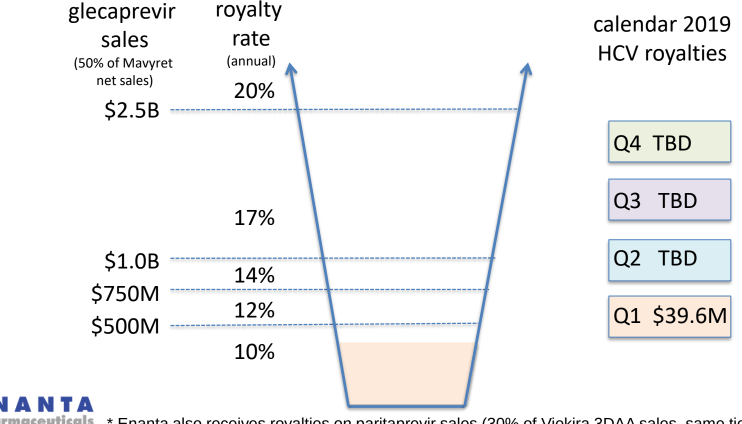




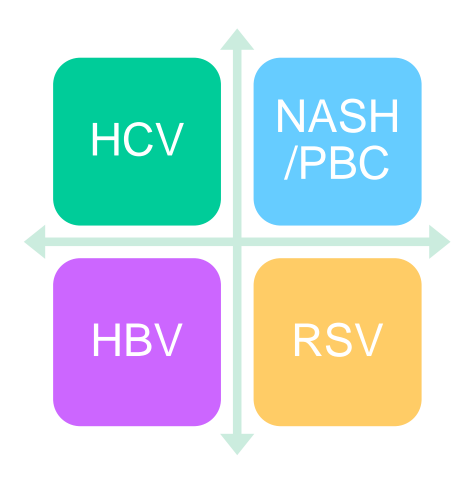


Glecaprevir The Pan-genotypic HCV Protease Inhibitor in AbbVie's MAVYRET™

Product	Regimen	Enanta Asset	Economics*
MAVYRET. glecaprevir/pibrentasvir	2-DAA (ABBV)	glecaprevir (PI)	Double-digit royalty on 50% of net sales



Virology & Liver Disease Focus Areas





Respiratory Syncytial Virus (RSV)

- Negative-sense, single-stranded RNA virus of family Pneumoviridae
- Can cause severe lung infections, including bronchiolitis (infection of small airways in the lungs) and pneumonia (an infection of the lungs)
- Higher risk populations for severe illness include:
 - Premature babies
 - Older adults, especially those 65 years and older
 - People with chronic lung disease or certain heart problems
 - People with weakened immune systems (e.g. HIV, organ transplant, chemotherapy)
- Each year in U.S.:
 - > 57,000 children below age 5 are hospitalized for RSV
 - ~ 177,000 older adults are hospitalized, and about 14,000 die
- No safe and effective treatments



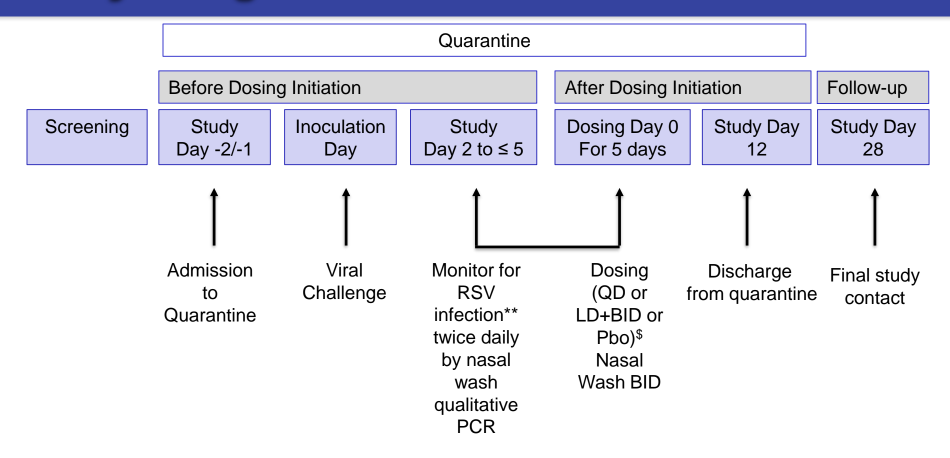
Source: CDC

EDP-938: Enanta's First Clinical-Stage Compound for RSV

- EDP-938 is the only N-inhibitor under clinical evaluation
 - Non-Fusion approach directly targets viral replication
- Strong preclinical virological profile:
 - Nanomolar inhibitor of both RSV-A and RSV-B activity
 - Maintained antiviral potency across all clinical isolates tested
 - Demonstrated high-barrier to resistance in vitro
 - Synergy with other drug mechanisms (e.g. fusion and L inhibitors)
 - Active against resistant virus from other mechanisms
 - Robust in vivo efficacy data
- Phase 2a human challenge study complete



Phase 2a Challenge Study (EDP 938-101) Study Design and Procedures

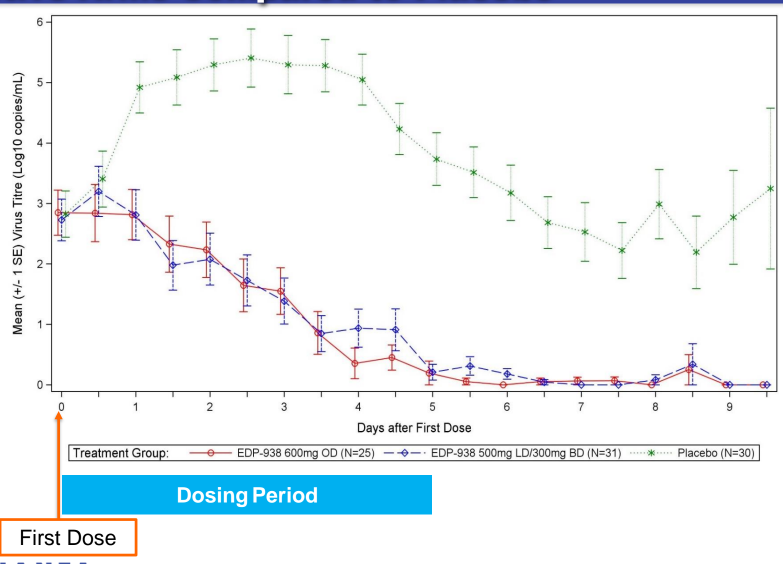


^{**} Dosing (D0) is initiated 12 hours after testing positive for RSV or Day 5 (PM), whichever comes first

- \$ EDP-938/placebo is administered as a blinded oral liquid suspension
- EDP-938 500mg loading dose, then 300mg BID over 5 days
- EDP-938 600mg QD Q24h alternating with placebo Q24h x 5 days to maintain the blind
- Placebo for EDP-938 BID x 5 days



Robust Antiviral Effect Rapid and Sustained Reduction in Viral Load in Both Active Arms Compared to Placebo



Highly Statistically Significant Reduction in Both EDP-938 Arms Compared to Placebo

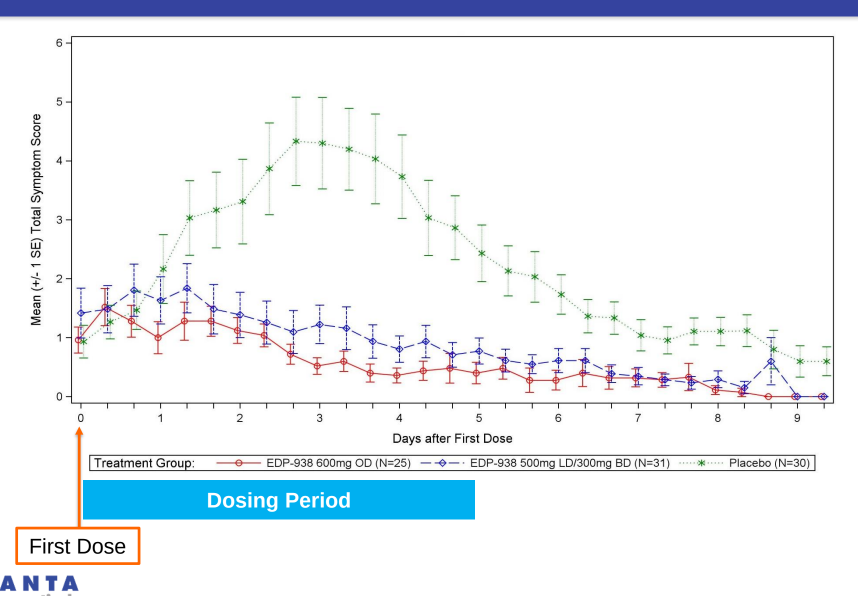
- No Statistically Significant Difference Between the Two Dosing Regimens

	EDP-938 600 mg QD	EDP-938 500 mg LD + 300 mg BID	Placebo
N	25	31	30
Viral load AUC mean (SD) (hours x Log ₁₀ copies/mL)	203.95 (173.50)	217.71 (217.55)	790.15 (408.80)
% Reduction (relative to placebo)	74.43%	71.46%	
Absolute Reduction* (relative to placebo)	-588.08	-564.63	
P-value	<0.001	<0.001	
Difference between two EDP-938 dosing groups	-23.45		
P-value	0.722		

^{*} Difference in LS Mean



EDP-938 Shows a Rapid and Sustained Attenuation of RSV Symptoms Compared to Placebo



Both EDP-938 Regimens Demonstrated Highly Statistically Significant Attenuation of RSV Symptoms Compared to Placebo - No Statistically Significant Difference Between the Two Dosing Regimens

	EDP-938 600 mg QD	EDP-938 500 mg LD/300 mg BID	Placebo
N	25	31	30
AUC Total Symptom Score mean (SD) (hours x Score)	124.47 (115.60)	181.75 (248.42)	478.75 (422.29)
% Reduction (relative to placebo)	74.3%	68.2%	
Absolute Reduction* (relative to placebo)	-355.91	-326.64	
P-value	<0.001	<0.001	
Difference between two EDP-938 dosing groups	-29.27		
P-value	0.700		

^{*} Difference in LS Mean



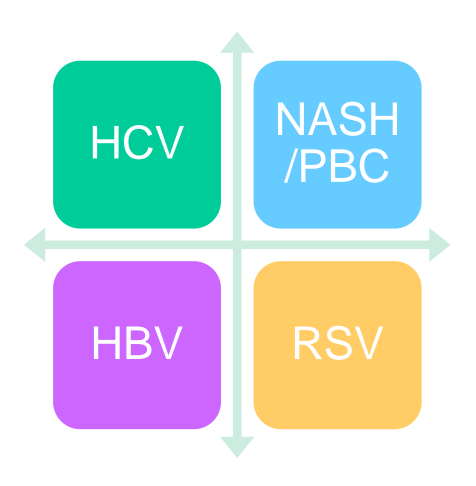
Summary: EDP-938, A Highly Efficacious and Safe RSV N-Inhibitor

Phase 1 results:

- Safe and well tolerate, no SAEs, AEs were mild
- At Phase 2 doses, mean trough levels 30x higher than EC 90 of EDP-938 against RSV-infected human cells
- Phase 2a Results:
 - Primary and Key Secondary Efficacy Endpoints were achieved (p<0.001) at both dose levels after 5 days of dosing
 - EDP-938 mean C_{trough} concentrations were approximately >20-40x higher than EC₉₀
 - Well tolerated with safety profiles similar to placebo (consistent profile observed in >250 subjects exposed to EDP-938 for up to 7 days in Phase 1 and Phase 2a)
- Future Phase 2 studies will focus on both adult and infant populations



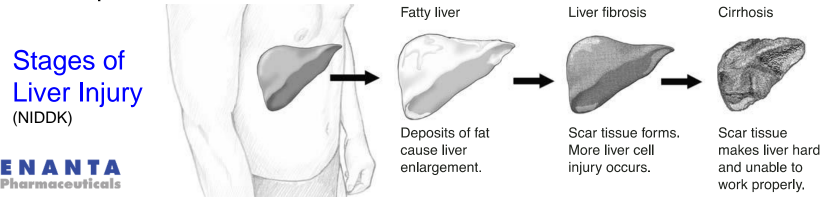
Virology & Liver Disease Focus Areas





Non-Alcoholic Fatty Liver Disease (NAFLD) and Non-Alcoholic Steatohepatitis (NASH)

- Number one cause of liver disease in Western Countries
- NAFLD: excessive fat (triglyceride) accumulation in the liver (steatosis)
- A subgroup of NAFLD patients has liver cell injury and inflammation in addition to excessive fat (steatohepatitis), i.e. NASH
- NASH is associated with the metabolic syndrome diseases related to type 2 diabetes, insulin resistance, obesity, hyperlipidemia, and hypertension
- While NAFLD does not correlate with short-term morbidity or mortality, but progression to NASH dramatically increases risks of cirrhosis, liver failure, and hepatocellular carcinoma



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Enanta's Approach to NASH and PBC-Agonists of Farnesoid X Receptor (FXR)

- FXR
 - nuclear receptor
 - main regulator of bile acid levels in liver and small intestine
 - responds to bile acids by regulating transcription of key enzymes and transporters

Inflammation

- FXR agonist preclinical PoC
 - ameliorate pathologies in NASH and PBC models, including an effect on fibrosis

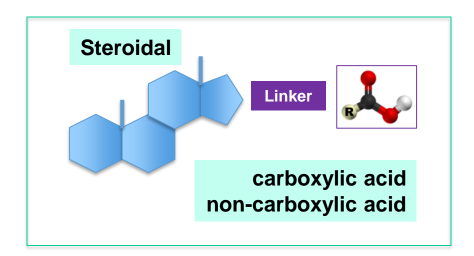
Bile Acid

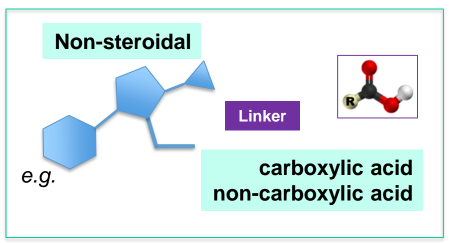
Regulation

 Clinical validation of FXR agonist in NASH and PBC with 6-ECDCA (OCA)



Classification of FXR Agonists – Four fundamental types (with variations)





FXR Agonists		Example
Steroidal carboxylic acid	S-CA	OCA, bile acids
Steroidal non-carboxylic acid	S-NCA	Enanta compounds
Non-steroidal carboxylic acid	NS-CA	Enanta compounds, GS-9674, LJN452
Non-steroidal non-carboxylic acid	NS-NCA	Enanta compounds



FXR Agonist EDP-305: Introduction

- EDP-305: Steroidal non-carboxylic acid, modified with additional non-steroidal binding element to enhance potency
- Potent FXR receptor agonist activity vs OCA
- Highly selective for FXR vs other nuclear receptors
 - and vs TGR5 receptor
- Potent and differentiated effects on FXR-dependent gene expression vs OCA
 - e.g. Shp, Cyp7a1, Bsep, Fgf15/FGF19
 - human hepatocytes and in vivo mouse model
- Efficacy in multiple NASH models
 - STAMTM mouse NASH model and dietary-induced NASH (DIN) mouse model
 - Improvement in hepatocyte ballooning and overall NAFLD Activity Score vs OCA
- Reduced liver fibrosis in rodent models
 - Mdr2-/-, MCD, CDAHFD, thioacetamide, and bile duct ligation models



EDP-305 Phase 1 Study

- Double-blind, placebo-controlled, Phase 1a/b study
- Healthy adults, and adults with presumptive NAFLD ("PN")
 - PN were obese, with or without pre-diabetes or type 2 diabetes mellitus, mean BMI= 32
- Oral suspension EDP-305 or placebo, dosed once daily
 - Total N=146 subjects (n=110 EDP305, n=36 pbo)
 - SAD, n=50, 6 cohorts at 1, 5, 10, 20, 40 and 80 mg
 - MAD, n=48 healthy and n=48 PN, 6 cohorts at 0.5, 1, 2.5, 5, 10, and 20 mg for 14 days
- Safety, tolerability, PK, and proof of target engagement support progression to Ph2 with once daily dosing

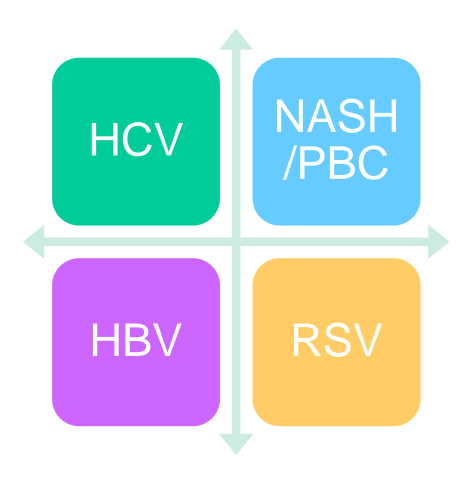


FXR Agonist EDP-305: Ph2 Studies

- Fast Track Designation granted by FDA for PBC and for NASH with fibrosis
- Two Ph 2 studies ongoing:
 - "ARGON-1" (NASH) and "INTREPID" (PBC)
 - 12 week dose ranging, randomized, double-blind, placebo-controlled
 - Evaluate safety, tolerability, PK, and efficacy (ALP reduction in PBC and ALT reduction in NASH)
 - New tablet formulation at 1 and 2.5 mg (~2X greater exposure than Ph1 suspension formulation)
 - ARGON-1 enrollment complete



Virology & Liver Disease Focus Areas





HBV Background

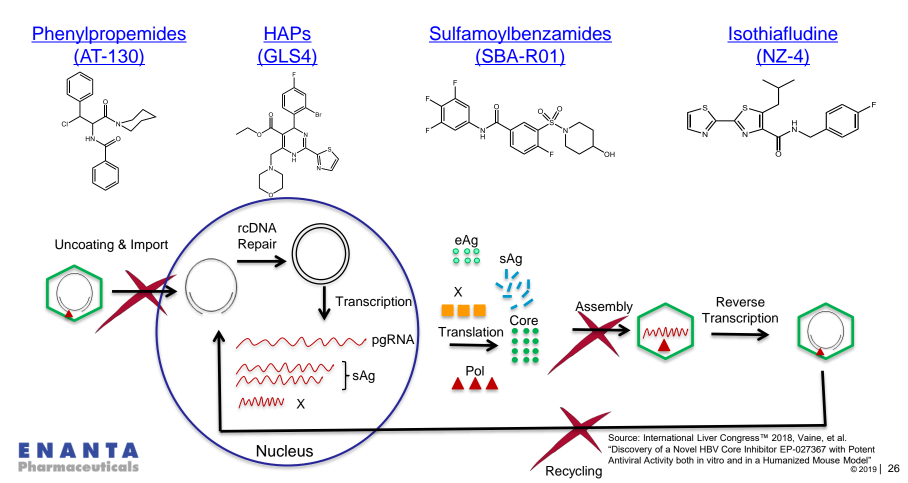
- Potentially life-threatening liver infection caused by the hepatitis B virus
- Current treatments rarely give true cures
 - Interferon gives better results (~10%), but with side effects
 - RT inhibitors very effective at reducing viral load, but offer very low cure rates (1% or lower) and must be taken for life to improve cirrhosis or HCC outcomes
- Prevalence estimates
 - US: ~850,000 2 million
 - US + Japan + major EU populations: ~4.9 million
 - Worldwide: ~250 million
- Estimated 15-25% of patients with chronic HBV infection will develop chronic liver diseases including cirrhosis, HCC, or liver decompensation



Core inhibitors: Introduction

(also called capsid assembly modulators, core protein allosteric modulators, capsid inhibitors)

- Novel class of replication inhibitor
- Act at multiple steps in HBV lifecycle
 - prevent proper uncoating, nuclear import, assembly, and recycling

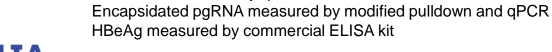


Core Inhibitor EDP-514 is a Potent Inhibitor of HBV Replication

EDP-514 is active in multiple HBV stable cell lines

	HBV Stable Cell Line EC ₅₀ (nM)			
	HepAD38	HepDE19	HepG2.2.15	
Intracellular Viral DNA	18	27	17	
Encapsidated pgRNA	25	3	5	
HBeAg	20	34	>500*	

^{*} In HepG2.2.15 cells, HBeAg is transcribed off transgene and is not dependent on viral replication



Viral DNA measured by qPCR



EDP-514 Prevents *de novo* Formation of cccDNA in Primary Human Hepatocytes

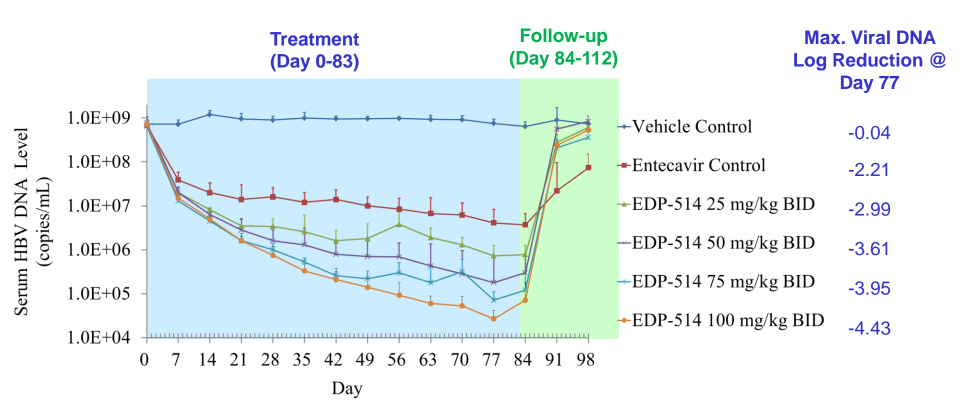
 EDP-514 prevents cccDNA establishment when present at early time points in infection (HBsAg as surrogate marker)

Compound	HBsAg EC ₅₀ (nM)		HBV DNA EC ₅₀ (nM)		
	d0 Addition	d3 Addition	d0 Addition	d3 Addition	
EDP-514	35	>1000	10	6	
Entecavir	>1000	>1000	0.25	0.21	



EDP-514 is Efficacious in the Humanized Liver Mouse Model

 uPA/SCID mice were infected with genotype C HBV and subsequently treated with EDP-514 BID at indicated doses for 12 weeks





HBV Core Inhibitor EDP-514 Summary

- A novel core inhibitor that displays potent anti-HBV activity at multiple points in the HBV lifecycle
- In vitro:
 - Potent anti-HBV activity in HBV expressing stable cells lines
 - Capable of preventing the establishment of cccDNA
 - Potent pan-genotypic activity
- In vivo:
 - Favorable tolerability and pharmacokinetic profile
 - Over 4-log reduction in HBV viral titers with 12 weeks of treatment in a chimeric liver mouse model
- Ph1 start targeted for 3Q-19



Financial Highlights

(\$ In millions)	Fiscal Year Ended Sept. 30, 2018	Fiscal 2Q19
Total Revenues	206.6*	\$39.6
R&D Expenses	\$94.9	\$34.2
G&A Expenses	\$23.4	\$6.8
Net Income	\$71.9	\$4.1
EPS (per diluted share)	\$3.48	\$0.20
Balance Sheet		
Cash, Cash Equivalents and Marketable Securities	\$325.1	\$386.7

^{*} Includes \$15M milestone payment from AbbVie for reimbursement approval of MAVIRET™ in Japan



Key Catalysts



- Ongoing double-digit HCV royalties from glecaprevir (MAVYRET™)
- RSV program:
 - Goal Initiate Ph2b adult outpatient study by end of 2019
- FXR agonist EDP-305 for NASH / PBC:
 - Phase 2 data in NASH by end of 3Q19
 - Identify follow-on FXR clinical candidate for NASH in 2019
 - Advance non-FXR compounds for NASH
 - Continued PBC enrollment in 2019
- HBV program
 - Initiate Phase 1 with Core Inhibitor EDP-514 in 3Q19



